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(54) Tide: PIPERIDINE DERIVATIVES USEFUL AS CCR5 ANTAGONISTS

NR*-C(0)-O-elkyl NR⁵-C(0)-NH-sikyl ਨੂੰ ਸੂ ਹੈ ਹੈ CH₂-ellsyl-R³ -CR¹³-NR⁵-C(O)-N-(alkyl)₂ -CR¹³-- (I

The use of CCR3 antagenists of formula (I) or a pharmaceutically acceptable salt thereof, wherein X is $-C(R^{13})_{R^{-1}}$, $-C(R^{13})$ (57) Abstract

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PIPERIDINE DERIVATIVES USEFUL AS CCR5 ANTAGONISTS

BACKGROUND

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The present invention relates to piperidine derivatives useful as selective CCR5 antagonists, pharmaceutical compositions containing the compounds, and methods of treatment using the compounds. The invention also relates to the use of a combination of a CCR5 antagonist of this invention and one or more antiviral or other agents useful in the treatment of Human Immunodeficiency Virus (HIV). The invention further relates to the use of a CCR-5 antagonist of this invention, alone or in combination with another agent, in the treatment of solid organ transplant rejection, graft v. host disease, arthritis, rheumatoid arthritis, inflammatory bowel disease, atopic dermatitis, psoriasis, asthma, allergies or multiple sclerosis.

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The global health crisis caused by HIV, the causative agent of Acquired Immunodeficiency Syndrome (AIDS), is unquestioned, and while recent advances in drug therapies have been successful in slowing the progression of AIDS, there is still a need to find a safer, more efficient, less expensive way to control the virus.

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It has been reported that the CCR5 gene plays a role in resistance to HIV infection. HIV infection begins by attachment of the virus to a target cell membrane through interaction with the cellular receptor CD4 and a secondary chemokine co-receptor molecule, and proceeds by replication and dissemination of infected cells through the blood and other tissue. There are various chemokine receptors, but for macrophage-tropic HIV, believed to be the key pathogenic strain that replicates *in vivo* in the early stages of infection, the principal chemokine receptor required for the entry of HIV into the cell is CCR5. Therefore, interfering with the interaction

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between the viral receptor CCR5. Therefore, interfering with the interaction between the viral receptor CCR5 and HIV can block HIV entry into the cell. The present invention relates to small molecules which are CCR5 antagonists.

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WO 00/66559

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CCR-5 receptors have been reported to mediate cell transfer in inflammatory diseases such as arthritis, rheumatoid arthritis, atopic dermatitis, psoriasis, asthma and allergies, and inhibitors of such receptors are expected to be useful in the treatment of such diseases, and in the treatment of other inflammatory diseases or conditions such as inflammatory bowel disease, multiple sclerosis, solid organ transplant rejection and graft v. host disease.

Related piperidine derivatives which are muscarinic antagonists useful in the treatment of cognitive disorders such as Alzheimer's disease are disclosed in US patents 5,883,096; 6,037,352; 5,889,006; 5,952,349; and 5,977,138.

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A-M. Vandamme et al., <u>Antiviral Chemistry & Chemotherapy</u>, 9:187-203 (1998) disclose current clinical treatments of HIV-1 infections in man including at least triple drug combinations or so-called Highly Active

15 Antiretroviral Therapy ("HAART"); HAART involves various combinations of nucleoside reverse transcriptase inhibitors ("NRTI"), non-nucleoside reverse transcriptase inhibitors ("NRTI"), non-nucleoside reverse transcriptase inhibitors ("NRTI") and HIV protease inhibitors ("PI"). In compliant drug-naive patients, HAART is effective in reducing mortality and progression of HIV-1 to AIDS. However, these multidrug therapies do not eliminate HIV-1 and long-term treatment usually results in multidrug resistance. Development of new drug therapies to provide better HIV-1 treatment remains a priority.

SUMMARY OF THE INVENTION

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The present invention relates to the treatment of HIV comprising administering to a human in need of such treatment an effective amount of a CCR5 antagonist represented by the structural formula I:

or a pharmaceutically acceptable salt thereof, wherein X is -C(A13)₂-, -C(A13)(A19)-, -C(O)-, -O-, -NH-, -N((C₁-C₆)alkyl)-,

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O-C(O)-(C₁-C₆)alkyl O-C(O)-O-(C₁-C₆)alkyl O-C(O)-NH-(C₁-C₆)alkyl O-C(O)-NH-(C₁-C₆)alkyl O-C(O)-NH-(C₁-C₆)alkyl

O-C(O)-N((C₁-C₆)alkyl)₂ NR⁵-C(O)-(C₁-C₆)alkyl -CR¹³- , -CR¹³-

NH⁵-C(O)-N-((C₁-C₆)alkyl)₂ C(O)-(C₁-C₆)alkyl

R is R6-phenyl, R6-pyridyl, R6-thiophenyl or R6-naphthyl; R1 is hydrogen, C1-C6 alkyl or C2-C6 alkenyl;

R² is R⁷, R⁸, R⁹-phenyl; R⁷, R⁸, R⁹-substituted 6-membered heteroaryl; R⁷, R⁸, R⁹-substituted 6-membered heteroaryl N-oxide; R¹⁰, R¹¹-substituted 5-membered heteroaryl; naphthyl; fluorenyl;

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HT R¹² RT C—heteroaryl

R3 is R6-phenyl, R*-heteroaryl or R*-naphthyl;

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- R⁴ is hydrogen, C₁-C₆ alkyl, fluoro-C₁-C₆ alkyl, cyclopropylmethyl, -CH₂CH₂OH, -CH₂CH₂-O-(C₁-C₆)alkyl, -CH₂C(O)-O-(C₁-C₆)alkyl, -CH₂C(O)-N((C₁-C₆)alkyl)₂; -CH₂C(O)NH₂, -CH₂C(O)-NH(C₁-C₆)alkyl or -CH₂C(O)-N((C₁-C₆)alkyl)₂; R⁵ and R¹¹ are independently selected from the group consisting of
- R⁶ is 1 to 3 substituents independently selected from the group consisting of hydrogen, halogen, C₁-C₆ alkyl, C₁-C₆ alkoxy, -CF₃, CF₃O-, CH₃C(O)-, -CN, CH₃SO₂-, CF₃SO₂-, R¹⁴-phenyl, R¹⁴-benzyl,

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hydrogen and (C1-C6)-alkyl;

$$\label{eq:ch3C} \begin{split} &\text{CH}_3\text{C}(=\text{NOCH}_3\text{C}(=\text{NOCH}_2\text{CH}_3)\text{-}, &\text{O}^{-1}\text{SO}_2^{-1}\text{-}\text{NH}_2, -\text{NHCOCF}_3,}\\ &\text{-NHCONH}(\text{C}_1\text{-C}_6\text{ alky}), -\text{NHCO}(\text{C}_1\text{-C}_6\text{ alky}), -\text{NHSO}_2(\text{C}_1\text{-C}_6\text{ alky}), \\ \end{split}$$

25 5-membered heteroaryl and , wherein X is -O-, -NH- or -N(CH₃)-;
R7 and R8 are independently selected from the group consisting of
(C₁-C₆)alkyl, halogen, -NR[∞]R³', -OH, -CF₃, -OCH₃, -O-acyl, and -OCF₃;
R8 is R7 budgeon should NC CN CU F Cuts

R⁹ is R⁷, hydrogen, phenyl, -NO₂, -CN, -CH₂F, -CHF₂, -CHO, -CH=NOR²⁰, pyridyl, pyridyl N-oxide, pyrimidinyl, pyrazinyl,

-N(R²⁶)CONR²¹R²², -NHCONH(chloro-(C₁-C₆)alkyl), -NHCONH((C₃-C₁₀)-

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WO 00/66559

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PCT/US00/11633

cycloalkyl(C₁-C₆)alkyl), -NHCO(C₁-C₆)alkyl, -NHCOCE₃, -NHSO₂N((C₁-C₆)alkyl)₂, -NHSO₂(C₁-C₆)alkyl, -N(SO₂CE₃)₂, -NHCO₂(C₁-C₆)alkyl, C₃-C₁₀ cycloalkyl, -SR²³, -SOR²³, -SO₂R²³, -SO₂NH(C₁-C₆ alkyl), -OSO₂(C₁-C₆)alkyl, -OSO₂CE₃, hydroxy(C₁-C₆)alkyl, -CON R²⁰R²³, -CON(CH₂CH₂-O-CH₃)₂,

-OCONH(C₁-C₆)alkyl, - \dot{C} O₂R²⁰, -Si(CH₃)₃ or -B(OC(CH₃)₂)₂: R¹⁰ is (C₁-C₆)alkyl, -NH₂ or R¹²-phenyl;

R¹² is 1 to 3 substituents independently selected from the group consisting of hydrogen, (C₁-C₆) alkyl, -CF₃, -CO₂R₂₀, -CN, (C₁-C₆)alkoxy

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R13, R14, R15 and R16 are independently selected from the group consisting of hydrogen and (C_1 - C_6)alkyl;

R¹⁷ and R¹⁸ are independently selected from the group consisting of hydrogen and C₁-C₆ alkyl, or R¹⁷ and R¹⁸ together are a C₂-C₅ alkylene group and with the carbon to which they are attached form a spiro ring of 3

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to 6 carbon atoms; R¹⁸ is R^e-phenyl, R^e-heteroaryl, R^e-naphthyl, C₃-C₁₀ cycloalkyl, (C₃-

 C_{10})cycloalkyl(C_1 - C_6)alkyl or (C_1 - C_6)alkoxy(C_1 - C_6)alkyl; R^{20} , R^{21} and R^{22} are independently selected from the group

20 consisting of H and C₁-C₆ alkyl; and R²³ is C₁-C₆ alkyl or phenyl.

Preferred are compounds of formula I wherein R is R⁶-phenyl, especially wherein R⁶ is a single substituent, and especially wherein the R⁶ substituent is in the 4-position. Also preferred are compounds of formula I wherein R¹³, R¹⁴, R¹⁵ and R¹⁶ are each hydrogen or methyl, especially hydrogen. Also preferred are compounds of formula I wherein X is -CHOR³, -C(R¹³)(R¹⁸)- or -C(=NOR⁴)-: a preferred definition for R³ is

-CHOR3, -C(R¹3)(R¹²)- or -C(=NOR⁴)-; a preferred definition for R³ is pyridyl, especially 2-pyridyl, a preferred definition for R⁴ is (C₁-C₆)alkyl, 30 especially methyl, ethyl or isopropyl, a preferred definition for R¹³ is hydrogen, and a preferred definition for R¹³ is R⁵-phenyl. For compounds of formula I, R¹ is preferably (C₁-C₆)alkyl, especially methyl.

In compounds of formula I, R² is preferably R⁷, R⁸, R⁹-phenyl, R⁷, R⁸, R⁹-pyridyl or an N-oxide thereof, or R⁷, R⁸, R⁹-pyrimidyl. When R² is pyridyl, it is preferably 3- or 4-pyridyl, and when pyrimidyl, it is preferably 5-pyrimidyl. The R⁷ and R⁸ substituents are preferably attached to carbon ring members adjacent to the carbon joining the ring to the rest of the molecule and the R⁹ substituent can be attached to any of the remaining

unsubstituted carbon ring members, for example as shown in the following

methyl; halogen, especially chloro; and -NH2. A preferred R9 substituent Preferred R7 and R8 substituents are: (C1-C6)alkyl, especially

by the structural formula I Also claimed are novel CCR5 antagonist compounds represented

ö or a pharmaceutically acceptable salt thereof, wherein (1) X^a is -C(R¹³)₂-, -C(R¹³)(R¹⁹)-, -C(O)-, -O-, -NH-, -N((C₁-C₆)alkyl)-

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NR⁵-C(O)-O-(C₁-C₆)alkyl NR⁵-C(O)-NH-(C₁-C₆)alkyl -CR¹³- -CR¹³-

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NR⁵-C(O)-N-((C₁-C₆)alkyl)₂ C(O)-(C₁-C₆)alkyl -CH¹³— or -N- ;

R1 is hydrogen, C1-C6 alkyl or C2-C6 alkenyl; Ra is Rea-phenyl, Rea-pyridyl, Rea-thiophenyl or Re-naphthy!

8R10, R11-substituted 5-membered heteroaryl; naphthyl; fluorenyl; heteroaryl; R7, R8, R9-substituted 6-membered heteroaryl N-oxide R² is R⁷, R⁸, R⁹-phenyl; R⁷, R⁸, R⁹-substituted 6-membered

PCT/US00/11633

diphenylmethyl

R⁴ is hydrogen, C₁-C₆ alkyl, fluoro-C₁-C₆ alkyl, cyclopropylmethyl, R3 is R10-phenyl, pyridyl, pyrimidyl, pyrazinyl or thiazolyl,

hydrogen and (C₁-C₆)-alkyl; -CH₂C(O)NH₂, -CH₂C(O)-NH(C₁-C₆)alkyl or -CH₂C(O)-N((C₁-C₆)alkyl)₂; --·CH₂CH₂OH, -CH₂CH₂-O-(C₁-C₆)alkyl, -CH₂C(O)-O-(C₁-C₆)alkyl, R5 and R11 are independently selected from the group consisting of

consisting of hydrogen, halogen, -CF3, CF3O-, -CN, -CF3SO2-, R12-phenyl, R^{6a} is 1 to 3 substituents independently selected from the group

ö -NHCOCF₃, 5-membered heteroaryl and , wherein X is -O-, -NH-

R6 is independently selected from the group consisting of R6a and

8 5 cycloalkyl, $-SR^{23}$, $-SOR^{23}$, $-SO_2R^{23}$, $-SO_2NH(C_1-C_6$ alkyl), $-OSO_2(C_1-C_6)$ alkyl, $-OSO_2CF_3$, hydroxy(C_1-C_6) alkyl, -CON $R^{20}R^{21}$, $-CON(CH_2CH_2-O-C_6)$ alkyl, -CON(C1-C6)alkyl, halogen, -NR²⁰R²¹, -OH, -CF3, -OCH3, -O-acyl, and -OCF3; C_6)alkyl)₂, -NHSO₂(C_1 - C_6)alkyl, -N(SO₂CF₃)₂, -NHCO₂(C_1 - C_8)alkyl, C_3 - C_{10} cycloalkyl(C₁-C₆)alkyl), -NHCO(C₁-C₆)alkyl, -NHCOCF₃, -NHSO₂N((C₁- $-N(R^{20})CONR^{21}R^{22}$, $-NHCONH(chloro-(C_1-C_6)aikyl)$, $-NHCONH((C_3-C_{10})-C_{10})$ -CH=NOR²⁰, pyridyl, pyridyl N-oxide, pyrimidinyl, pyrazinyl, R9 is R', hydrogen, phenyl, -NO₂, -CN, -CH₂F, -CHF₂, -CHO, R7 and R8 are independently selected from the group consisting of

25 -OCONH(C_1 - C_6)alkyl, - CO_2R^{20} , -Si(CH_3)₃ or -B($OC(CH_3)_2$)₂ R¹⁰ is (C₁-C₆)alkyl, -NH₂ or R¹²-phenyl;

consisting of hydrogen, (C₁-C₆) alkyl, -CF₃, -CO₂R₂₀, -CN, (C₁-C₆)alkoxy R¹² is1 to 3 substituents independently selected from the group

မ consisting of hydrogen and (C₁-C₆)alkyl; R13, R14, R15 and R16 are independently selected from the group

group and with the carbon to which they are attached form a spiro ring of 3 hydrogen and C_1 - C_6 alkyl, or R¹⁷ and R¹⁸ together are a C_2 - C_5 alkylene R¹⁷ and R¹⁸ are independently selected from the group consisting of

WO 00/66559

PCT/US00/11633

C₁₀)cycloalkyl(C₁-C₆)alkyl or (C₁-C₆)alkoxy(C₁-C₆)alkyl; R²⁰, R²¹ and R²² are independently selected from the group R" is R°-phenyl, R°-heteroaryl, R°-naphthyl, C3-C10 cycloalkyl, (C3-

R²³ is C₁-C₆ alkyl or phenyl; or

consisting of H and C1-C8 alkyl; and

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Xª is -C(R13)(R19)-, -C(O)-, -O-, -NH-, -N((C1-C6)alkyl)-

O-C(O)-(C1-C6)alkyl O-C(O)-NH-(C1-C6)alkyl CR13- - CR13-

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NR5-C(O)-N-((C1-C6)alkyl)2 C(O)-(C1-C6)alkyl Ra is R6b-phenyl, R6b-pyridyl or R6b-thiophenyl;

8 $- CH_2CH_2 - O - (C_1 - C_6) \\ alkyl, - CH_2C(O) - O - (C_1 - C_6) \\ alkyl, - CH_2C(O) \\ NH_2, - CH_2C(O) \\ - CH_2CH_2 - CH_2CH_2 - CH_2CH_2 \\ - CH_2CH_2 - CH_2CH_2 - CH_2 \\ - CH_2CH_2 - CH_2 - CH_2 \\ - CH_2CH_2 - CH_2 - CH_2 \\ - CH_2 - CH_2 - CH_2 - CH_2 - CH_2 \\ - CH_2 - CH_2 - CH_2 - CH_2 - CH_2 \\ - CH_2 - CH_2 - CH_2 - CH_2 - CH_2 \\ - CH_2 - CH_2 - CH_2 - CH_2 - CH_2 \\ - CH_2 \\ - CH_2 - CH_2$ CH2C(O)-NH-(C1-C6)alkyl or -CH2C(O)-N((C1-C6)alkyl)2; R6b is CH3SO2-; and R^{4a} is fluoro-C₁-C₆ alkyl, cyclopropylmethyl, -CH₂CH₂OH

R1, R2, R3, R5, R14, R15, R16 and R19 are as defined in (1)

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definition for \mathbb{R}^4 is $(C_1\text{-}C_6)$ alkyl, especially methyl, ethyl or isopropyl, a Rephenyl. For compounds of formula II(1), R1 is preferably (C1-C6)alkyl preferred definition for R¹³ is hydrogen, and a preferred definition for R¹⁹ is especially wherein \mathbf{R}^{6a} is a single substituent, and especially wherein the preferred definition for R3 is pyridyl, especially 2-pyridyl, a preferred R^{6a} substituent is in the 4-position. Also preferred are compounds of formula II(1) wherein X^a is -CHOR3, -C(R¹³)(R¹⁸)- or -C(=NOR⁴)-; a Preferred are compounds of formula II(1) wherein Ra is R6a-pheny!

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are preferably hydrogen. especially methyl. Also for compounds of formula II(1), R14, R15 and R16

preferred definition for R3 is pyridyl, especially 2-pyridyl, preferred R6b substituent is in the 4-position. Also preferred are compounds of definition for R13 is hydrogen, and a preferred definition for R18 is R5-phenyl. definitions for R^{4a} are cyclopropylmethyl and trifluoroethyl, a preferred formula II(2) wherein Xa is -CHOR3, -C(R13)(R19)- or -C(=NOR4a)-; a especially wherein R6b is a single substituent, and especially wherein the Preferred are compounds of formula II(2) wherein Ra is R6b-phenyl

preferably hydrogen. In compounds of formula II(1) and (2), R2 is preferably R7, R8, R9.

methyl. Also for compounds of formula II(2), R14, R15 and R16 are For compounds of formula II(2), R1 is preferably (C1-C6)alkyl, especially

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20 ជ compounds of formula 1. Preferred R7 and R8 substituents for compounds chloro; and -NH2; a preferred R9 substituent is hydrogen. When R² is pyridyl, it is preferably 3- or 4-pyridyl, and when pyrimidyl, it is of formula II are: (C1-C6)alkyl, especially methyl; halogen, especially the molecule and the R9 substituent can be attached to any of the to carbon ring members adjacent to the carbon joining the ring to the rest of preferably 5-pyrimidyl. The R7 and R8 substituents are preferably attached phenyl; R7, R8, R9-pyridyl or an N-oxide thereof; or R7, R8, R9-pyrimidyl. remaining unsubstituted carbon ring members as shown above for

treatment of HIV comprising an effective amount of a CCR5 antagonist of Another aspect of the invention is a pharmaceutical composition for

မွ 23 amount of a CCR5 antagonist of formula II in combination with a treatment of solid organ transplant rejection, graft v. host disease, arthritis, psoriasis, asthma, allergies or multiple sclerosis comprising an effective Another aspect of the invention is a pharmaceutical composition for formula II in combination with a pharmaceutically acceptable carrier. rheumatoid arthritis, inflammatory bowel disease, atopic dermatitis,

comprising administering to a human in need of such treatment an effective pharmaceutically acceptable carrier. Yet another aspect of this invention is a method of treatment of HIV

႘ၟ graft v. host disease, arthritis, rheumatoid arthritis, inflammatory bowel the invention is a method of treatment of solid organ transplant rejection, disease, atopic dermatitis, psoriasis, asthma, allergies or multiple sclerosis amount of a CCR5 antagonist compound of formula II. Another aspect of

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amount of a CCR5 antagonist compound of formula I or II. comprising administering to a human in need of such treatment an effective

ഗ a CCR5 antagonist of formula I or II of this invention in combination with for the treatment of AIDS. Still another aspect of this invention is the use of or other agents useful in the treatment of Human Immunodeficiency Virus of formula I or II of this invention in combination with one or more antiviral one or more other agents useful in the treatment of solid organ transplant rejection, graft v. host disease, inflammatory bowel disease, rheumatoid Still another aspect of this invention is the use of a CCR5 antagonis

5 are components of the combination can be administered in a single dosage arthritis or multiple sclerosis. The CCR5 and antiviral or other agents which dosage forms of the actives is also contemplated. form or they can be administered separately; a kit comprising separate

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5 DETAILED DESCRIPTION OF THE INVENTION

unless otherwise indicated. As used herein, the following terms are used as defined below

from one to six carbon atoms. dialkylamino) represents straight and branched carbon chains and contains Alkyl (including the alkyl portions of alkoxy, alkylamino and

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unsaturated bonds, provided that two unsaturated bonds are not adjacent Alkenyl represents C2-C6 carbon chains having one or two

at any available position on the phenyl ring. Substituted phenyl means that the phenyl group can be substituted

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aralkyl is aryl-(C1-C6)alkyl, wherein aryl is as defined above. C7)cycloalkyl-(C1-C6)alkyl-C(O)-, and heteroaryl-C(O)-, wherein alkyl and alkyl-C(O)-, aryl-C(O)-, aralkyl-C(O)-, (C₃-C₇)cycloalkyl-C(O)-, (C₃heteroaryl are as defined herein; aryl is R12-phenyl or R12-naphthyl; and Acyl means a radical of a carboxylic acid having the formula

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structure and having a sufficient number of delocalized pi electrons to selected from O, S or N, said heteroatom(s) interrupting a carbocyclic ring bicyclic groups of 11 to 12 atoms having 1 or 2 heteroatoms independently Heteroaryl represents cyclic aromatic groups of 5 or 6 atoms or

႘ၟ provide aromatic character, provided that the rings do not contain adjacent form an N-oxide. All regioisomers are contemplated, e.g., 2-pyridyl, 3atoms can be substituted by R⁷, R⁸ or R⁹ groups. Nitrogen atoms can oxygen and/or sulfur atoms. For 6-membered heteroaryt rings, carbon

> WO 00/66559 PCT/US00/11633

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quinazolinyl, benzofuranyl, benzothienyl and indolyl. 4-position. Bicyclic groups typically are benzo-fused ring systems derived membered rings having two heteroatoms are preferably joined through the thiazolyl, isothiazolyl, imidazolyl, pyrazolyl and isoxazolyl. 5-Membered groups. Typical 5-membered heteroaryl rings are furyl, thienyl, pyrrolyl, membered heteroaryl rings, carbon atoms can be substituted by R10 or R11 pyrimidinyl, pyrazinyl, pyridazinyl and the N-oxides thereof. For 5from the heteroaryl groups named above, e.g. quinolyl, phthalazinyl, rings having one heteroatom can be joined through the 2- or 3- position; 5pyridyl and 4-pyridyl. Typical 6-membered heteroaryl groups are pyridyl, …

ᆳ preferably attached to a carbon ring member adjacent to the carbon joining is preferably alkyl; however, if a heteroatom is adjacent to the carbon ${\sf R}^2$ are described above. When ${\sf R}^2$ is a 5-membered heteroaryl group, the joining the ring to the rest of the molecule (i.e., as in 2-pyrrolyl), R10 is adjacent to the carbon joining the ring to the rest of the molecule, and R11 R¹⁰ and R¹¹ substituents are preferably attached to carbon ring members Preferred points of substitution for 6-membered heteroaryl rings at

the ring to the rest of the molecule.

Halogen represents fluoro, chloro, bromo and iodo.

20 different carbon atoms, e.g., -CH₂F, -CHF₂, -CF₃, F₃CCH₂- and -CF₂CF₃. substituted by 1 to 5 fluoro atoms, which can be attached to the same or Fluoro(C₁-C₆)alkyl represents a straight or branched alkyl chain

amount sufficient to lower HIV-1-RNA plasma levels. A therapeutically effective amount of a CCR5 antagonist is an

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antiviral agent or agents may be administered simultaneously or CCR5 antagonist in a single dosage form, or the CCR5 antagonist and the present invention. The antiviral agent or agents may be combined with the HIV-1 therapy may be used in combination with a CCR5 antagonist of the One or more, preferaby one to four, antiviral agents useful in anti-

မ္ဟ ဗ particular, the combinations known as HAART are contemplated for use in antiviral drugs listed below not falling within these classifications. In nucleoside reverse transcriptase inhibitors, protease inhibitors and other comprise nucleoside and nucleotide reverse transcriptase inhibitors, nonfor use in combination with the compounds of the present invention sequentially as separate dosage forms. The antiviral agents contemplated

("NRTI" s) as used herein means nucleosides and nucleotides and The term "nucleoside and nucleotide reverse transcriptase inhibitors"

combination with the CCR5 antagonists of this invention

-11-

analogues thereof that inhibit the activity of HIV-1 reverse transcriptase, the enzyme which catalyzes the conversion of viral genomic HIV-1 RNA into proviral HIV-1 DNA.

Typical suitable NRTIs include zidovudine (AZT) available under the RETROVIR tradename from Glaxo-Wellcome Inc., Research Triangle, NC 27709; didanosine (ddl) available under the VIDEX tradename from Bristol-Myers Squlbb Co., Princeton, NJ 08543; zalcitabine (ddC) available under the HIVID tradename from Roche Pharmaceuticals, Nutley, NJ 07110; stavudine (d4T) available under the ZERIT trademark from Bristol-

10 Myers Squibb Co., Princeton, NJ 08543; lamivudine (3TC) available under the EPIVIR tradename from Glaxo-Wellcome Research Triangle, NC 27709; abacavir (1592U89) disclosed in WO96/30025 and available under the ZIAGEN trademark from Glaxo-Wellcome Research Triangle, NC 27709; adefovir dipivoxil [bis(POM)-PMEA] available under the PREVON

tradename from Gilead Sciences, Foster City, CA 94404; lobucavir (BMS-180194), a nucleoside reverse transcriptase inhibitor disclosed in EP-0358154 and EP-0736533 and under development by Bristol-Myers Squibb, Princeton, NJ 08543; BCH-10652, a reverse transcriptase inhibitor (in the form of a racemic mixture of BCH-10618 and BCH-10619) under

development by Biochem Pharma, Laval, Quebec H7V, 4A7, Canada; emitricitabine [(-)-FTC] licensed from Emory University under Emory Univ. U.S. Patent No. 5,814,639 and under development by Triangle Pharmaceuticals, Durham, NC 27707; beta-L-FD4 (also called beta-L-D4C and named beta-L-2', 3'-dicleoxy-5-fluoro-cytidene) licensed by Yale

25 University to Vion Pharmaceuticals, New Haven CT 06511; DAPD, the purine nucleoside, (-)-beta-D-2,6,-diamino-purine dioxolane disclosed in EP 0656778 and licensed by Emory University and the University of Georgia to Triangle Pharmaceuticals, Durham, NC 27707; and lodenosine (FddA), 9-(2,3-dideoxy-2-fluoro-b-D-threo-pentofuranosyl)adenine, an acid stable

The term "non-nucleoside reverse transcriptase inhibitors" ("NNRTI"s) as used herein means non-nucleosides that inhibit the activity of HIV-1 reverse transcriptase.

purine-based reverse transcriptase inhibitor discovered by the NIH and under development by U.S. Bioscience Inc., West Conshohoken, PA

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Typical suitable NNRTIs include nevirapine (BI-RG-587) available under the VIRAMUNE tradename from Boehringer Ingelheim, the manufacturer for Roxane Laboratories, Columbus, OH 43216; delaviradine

WO 00/66559 PCT/US00/11633

- 12

(BHAP, U-90152) available under the RESCRIPTOR tradename from Pharmacia & Upjohn Co., Bridgewater NJ 08807; efavirenz (DMP-266) a benzoxazin-2-one disclosed in WO94/03440 and available under the SUSTIVA tradename from DuPont Pharmaceutical Co., Wilmington, DE 19880-0723; PNU-142721, a furopyridine-thio-pyrimide under development by Pharmacia and Upjohn, Bridgewater NJ 08807; AG-1549 (formerly Shionogi # S-1153); 5-(3,5-dichlorophenyl)- thio-4-isopropyl-1-(4-pyridyl)methyl-IH-imidazol-2-ylmethyl carbonate disclosed in WO 96 /10019 and under clinical development by Agouron Pharmaceuticals, Inc., LaJolla CA 92037-1020; MKC-442 (1-(ethoxy-methyl)-5-(1-methylethyl)-6-

(phenylmethyl)-(2,4(1H,3H)-pyrimidinedione) discovered by Mitsubishi Chemical Co. and under development by Triangle Pharmaceuticals, Durham, NC 27707; and (+)-calanolide A (NSC-675451) and B, coumann derivatives disclosed in NIH U.S. Patent No. 5,489,697, licensed to Med 15 Chem Research, which is co-developing (+) calanolide A with Vita-Invest as an orally administrable product.

The term "protease inhibitor" ("P") as used herein means inhibitors of the HIV-1 protease, an enzyme required for the proteolytic cleavage of viral polyprotein precursors (e.g., viral GAG and GAG Pol polyproteins), to into the individual functional proteins found in infectious HIV-1. HIV

20 into the individual functional proteins found in infectious HIV-1. HIV protease inhibitors include compounds having a peptidomimetic structure, high molecular weight (7600 daltons) and substantial peptide character, e.g. CRIXIVAN(available from Merck) as well as nonpeptide protease inhibitors e.g., VIRACEPT (available from Agouron).

Typical suitable PIs include saquinavir (Ro 31-8959) available in hard gel capsules under the INVIRASE tradename and as soft gel capsules under the FORTOVASE tradename from Roche Pharmaceuticals, Nutley, NJ 07110-1199; ritonavir (ABT-538) available under the NORVIR tradename from Abbott Laboratories, Abbott Park, IL 60064; indinavir (MK-30) available under the CRIXIVAN tradename from Merck & Co., Inc.,

30 639) available under the CRIXIVAN tradename from Merck & Co., Inc., West Point, PA 19486-0004; nelfnavir (AG-1343) available under the VIRACEPT tradename from Agouron Pharmaceuticals, Inc., LaJolla CA 92037-1020; amprenavir (141W94), tradename AGENERASE, a nonpeptide protease inhibitor under development by Vertex Pharmaceuticals, Inc., Cambridge, MA 02139-4211 and available from Glaxo-Wellcome,

Inc., Cambridge, MA 02139-4211 and available from Glaxo-Wellcome, Research Triangle, NC under an expanded access program; lasinavir (BMS-234475) available from Bristol-Myers Squibb, Princeton, NJ 08543 (originally discovered by Novartis, Basel, Switzerland (CGP-61755); DMP-

WO 00/66559 PCT/US00/11633

Pharmaceuticals, Inc., LaJolla CA 92037-1020. by Shionogi (Shionogi #S-1153) and under development by Agouron IL 60064; and AG-1549 an orally active imidazole carbamate discovered generation HIV-1 PI; ABT-378 under development by Abbott, Abbott Park development by Bristol-Myers Squibb, Princeton, NJ 08543, as a 2nd-Triangle Pharmaceuticals; BMS-2322623, an azapeptide under 450, a cyclic urea discovered by Dupont and under development by

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pentafuside and Yissum Project No. 11607. Hydroyurea (Droxia), a Other antiviral agents include hydroxyurea, ribavirin, IL-2, IL-12,

- ಠ by Bristol-Myers Squibb; In preclinical studies, it was shown to have a stavudine. IL-2 is disclosed in Ajinomoto EP-0142268, Takeda EPsynergistic effect on the activity of didanosine and has been studied with activation of T-cells, was discovered at the NCI and is under development ribonucleoside triphosphate reductase inhibitor, the enzyme involved in the
- ជ 0176299, and Chiron U. S. Patent Nos. RE 33653, 4530787, 4569790 94608-2997 as a lyophilized powder for IV infusion or sc administration PROLEUKIN (aldesleukin) tradename from Chiron Corp., Emeryville, CA 4604377, 4748234, 4752585, and 4949314, and is available under the upon reconstitution and dilution with water; a dose of about 1 to about 20
- 8 preferred. IL-12 is disclosed in WO96/25171 and is available from Roche million IU/day, sc is preferred; a dose of about 15 million IU/day, sc is more microgram/kg/day, sc is preferred. Pentafuside (DP-178, T-20) a 36-amino Madison, NJ 07940; a dose of about 0.5 microgram/kg/day to about 10 Pharmaceuticals, Nutley, NJ 07110-1199 and American Home Prodocts
- 25 Duke University to Trimeris which is developing pentafuside in collaboration acid synthetic peptide, disclosed in U.S. Patent No.5,464,933 licensed from membranes. Pentafuside (3-100 mg /day) is given as a continuous sc with Duke University; pentafuside acts by inhibiting fusion of HIV-1 to target infusion or injection together with efavirenz and 2 PI's to HIV-1 positive
- မ -1 VII protein, is under preclinical development by Yissum Research preferred. Yissum Project No. 11607, a synthetic protein based on the HIV patients refractory to a triple combination therapy; use of 100 mg/day is Development Co., Jerusalem 91042 , Israel. Ribavirin, 1-B-D-ribofuranosyl 1H-1,2,4-triazole-3-carboxamide, is available from ICN Pharmaceuticals,
- Inc., Costa Mesa, CA; its manufacture and formulation are described in U.S. Patent No. 4,211,771

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drug found useful for treating HIV-1 infections in man alone, or as part of The term "anti-HIV-1 therapy" as used herein means any anti-HIV-

WO 00/66559

PCT/US00/11633

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a second PI, and one NNRTI; and (ii) at least two anti-HIV-1 drugs selected quadruple combination therapies. Typical suitable known anti-HIV-1 such as (i) at least three anti-HIV-1 drugs selected from two NRTIs, one PI therapies include, but are not limited to multidrug combination therapies therapies include: from NNRTIs and PIs. Typical suitable HAART - multidrug combination multidrug combination therapies, especially the HAART triple and

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- 5 ಠ such as two NRTIs, one PI and a second PI or one NNRTI. In treatment of viral load plateau, a fourth drug,e.g., one PI or one NNRTI could be added combination therapy; the use of two NRTIs and one PI is prefered unless naive patients, it is preferred to start anti-HIV-1 treatment with the triple there is intolerance to PIs. Drug compliance is essential. The CD4⁺ and (b) two NRTIs and one NNRTI; and (c) quadruple combination therapies HIV-1-RNA plasma levels should be monitored every 3-6 months. Should (a) triple combination therapies such as two NRTIs and one PI; or
- See the table below wherein typical therapies are further described: ANTI-HIV-1 MULTI DRUG COMBINATION THERAPIES

Triple Combination Therapies

- Two NRTIs1 + one PI2
- 8 Two NRTIs1 + one NNRTI3

Quadruple Combination Therapies*

Two NRTIs + one PI + a second PI or one NNRTI

25 C. ALTERNATIVES:5

Two NRTI

One NRTI5 + one PI2

Two PIs⁶ ± one NRTI⁷ or NNRTI³

One PI2 + one NRTI7 + one NNRTI3

FOOTNOTES TO TABLE

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- One of the following: zidovudine + lamivudine; zidovudine + zidovudine + zalcitabine didanosine; stavudine + lamivudine; stavudine + didanosine;
- Indinavir, nelfinavir, ritonavir or saquinavir soft gel capsules.

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- Nevirapine or delavirdine.
- 9:187 at p 193-197 and Figures 1 + 2. See A-M. Vandamne et al Antiviral Chemistry & Chemotherapy

WO 00/66559

PCT/US00/11633

- 15 -

5. Alternative regimens are for patients unable to take a recommended regimen because of compliance problems or toxicity, and for those who fail or relapse on a recommended regimen. Double nucleoside combinations may lead to HIV-resistance and clinical failure in many patients.

Most data obtained with saquinavir and ritonavir (each 400 mg bid). Ċ,

- Zidovudine, stavudine or didanosine.
- Agents known in the treatment of rheumatoid arthritis, transplant and graft v. host disease, inflammatory bowel disease and multiple sclerosis which can be administered in combination with the CCR5 antagonists of the present invention are as follows:
- solid organ transplant rejection and graft v. host disease: immune
 15 suppressants such as cyclosporine and Interleukin-10 (IL-10), tacrolimus,
 antilymphocyte globulin, OKT-3 antibody, and steroids;

inflammatory bowel disease: IL-10 (see US 5,368,854), steroids and ulfidine;

rheumatoid arthritis: methotrexate, azathioprine, cyclophosphamide steroids and mycophenolate mofetil;

multiple sclerosis: interferon-beta, interferon-alpha, and steroids

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Certain CCR5 antagonist compounds of the invention may exist in different isomeric (e.g., enantiomers, diastereoisomers and atropisomers) forms. The invention contemplates all such isomers both in pure form and

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in admixture, including racemic mixtures.

Certain compounds will be acidic in nature, e.g. those compounds which possess a carboxyl or phenolic hydroxyl group. These compounds

which possess a carboxyl or phenolic hydroxyl group. These compounds may form pharmaceutically acceptable salts. Examples of such salts may include sodium, potassium, calcium, aluminum, gold and silver salts. Also contemplated are salts formed with pharmaceutically acceptable amines such as ammonia, alkyl amines, hydroxyalkylamines, N-methylglucamine

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Certain basic compounds also form pharmaceutically acceptable
35 salts, e.g., acid addition salts. For example, the pyrido-nitrogen atoms may
form salts with strong acid, while compounds having basic substituents
such as amino groups also form salts with weaker acids. Examples of
suitable acids for salt formation are hydrochloric, sulfuric, phosphoric,

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WO 00/66559 . PCT/US00/11633

-16-

acetic, citric, oxalic, malonic, salicylic, malic, fumaric, succinic, ascorbic, maleic, methanesulfonic and other mineral and carboxylic acids well known to those in the art. The salts are prepared by contacting the free base form with a sufficient amount of the desired acid to produce a salt in the

5 conventional manner. The free base forms may be regenerated by treating the salt with a suitable dilute aqueous base solution such as dilute aqueous NaOH, potassium carbonate, ammonia and sodium bicarbonate. The free base forms differ from their respective salt forms somewhat in certain physical properties, such as solubility in polar solvents, but the acid and base salts are otherwise equivalent to their respective free base forms for purposes of the invention.

All such acid and base salts are intended to be pharmaceutically acceptable salts within the scope of the invention and all acid and base salts are considered equivalent to the free forms of the corresponding compounds for purposes of the invention.

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Compounds of the invention can be made by the procedures known in the art, for example by the procedures described in the following reaction schemes, by the methods described in the examples below, and by using the methods described in US patents 5,883,096; 6,037,352; 5,889,006; 5,952,349; and 5,977,138.

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The following solvents and reagents may be referred to herein by the abbreviations indicated: tetrahydrofuran (THF); ethanol (EtOH); methanol (MeOH); acetic acid (HOAc or AcOH); ethyl acetate (EtOAc); N,N-dimethylformamide (DMF); trifluoroacetic acid (TFA); trifluoroacetic anhydride (TFAA); 1-hydroxy-benzotriazole (HOBT); m-chloroperbenzoic acid (MCPBA); triethylamine (EtgN); diethyl ether (EtgO); tert-butoxy-

anhydride (TFAA); 1-hydroxy-benzotriazole (HOBT); m-chloroperbenzoic acid (MCPBA); triethylamine (Et₃N); diethyl ether (Et₂O); tert-butoxy-carbonyl (BOC); 1,8-diazabicyclo[5.4.0]undec-7-ene (DBU); dimethyl-sulfoxide (DMSO); p-toluene sulfonic acid (p-TSA); potassium bis(trimethylsilyl)-amide (KHMDA); 4-dimethylaminopryidine (DMAP);
 N,N,N-diiospropylethylamine (Dipea); and 1-(3-dimethyl-aminopropyl)-3-ethyl carbodilimide hydrochloride (DEC). AT is room temperature.

ethyl carbodilmide hydrochloride (DEC). RT is room temperature.

Compounds of formula I and II wherein X is CHO(C=O)-(C₁-C₆)
alkyl, CHO(C=O)-(C₁-C₆)alkoxy, CHO(C=O)-NH-(C₁-C₆)alkyl,

CHNR⁵(C=O)-(C₁-C₆)alkyl, CHNR⁵(C=O)-(C₁-C₆)alkoxy, CHNR⁵(C=O)
CHNR⁵(C-O)-Ikyl or CHOR3 (and wherein B14 B15 and B16 are hydrocal).

35 NH-(C₁-C₆)alkyl or -CHOR³ (and wherein R¹⁴, R¹⁵ and R¹⁶ are hydrogen) are prepared according to Schemes 1-4:

WO 00/66559 PCT/US00/11633

- 17 -

Scheme '

4b. R3a = COR3 4a. R^{3a} = alkyl etc

prepared as depicted in Scheme 1. Ketone 1, the synthesis of which was described in WO98/05292, was subjected to standard amidation with formula I, Z is CH or N, and R 1 is an alkyl group such as methyl were Compounds of formula 3, wherein R, R7 and R8 are as defined for ഗ

- ō ArCOOH, EDCI or DEC, and HOBT, or ArCOCI, wherein Ar is R7, R8. Derivatization of the free hydroxyl moiety with alkyl halides, acyl chlorides substituted phenyl or pyridyl, followed by reduction with NaBH $_4$ to obtain 3. afforded ethers 4a, esters 4b, carbonates 4c, and carbamates 4d, (R3COCI), alkyl chloroformates (CICOOR3) and isocyanides (O=C=NR3)
- 5 were obtained after condensation of the hydroxyl 3 with phenyl or pyridyl respectively, wherein \mathbf{R}^3 is a lower alkyl group. The aryloxy compounds, $\mathbf{5}$ halides in the presence of a base.

Scheme 2

WO 00/66559

PCT/US00/11633

- 18 -

ಕ amide is performed as in Scheme 1. This route allows the introduction of various aryloxy and heteroaryloxy moieties at H3 through the use of C₆ lower alkyl. Removal of the Boc protecting group and conversion to the base as shown in Scheme 2, or by a hydroxy-substituted aryl or heteroaryl the free hydroxyl group with a halogen-substituted aryl in the presence of a of the N-Boc ketone 1a to the alcohol 6 first, followed by functionalization of nucleophilic displacement or Mitsunobu-type reaction on intermediate 6. azodicarboxylate of the formula R19O₂C-N=N-CO₂R²⁰, wherein R²⁰ is C₁-(wherein Z1 is as defined in Scheme 1) in the presence of PPh3 and an Alternatively, compounds of formula 5 can be prepared by reduction

5 oxime group with CH3ONH2·HCl, and reduction with BH3·S(CH3)2 to described in Scheme 1, were prepared by conversion of the ketone 2 to an Compounds of formula 8, wherein R, R1, R7, R8 and Z are as

- 19 -

chloroformate (CICOOR 20 , wherein R 20 is C1-C6 alkyl) or an isocyanide provide amine 8. Derivatization of the free amine moiety with an alkyl respectively. (O=C=NR3) affords carbamate compounds 9 and urea compounds 10,

Scheme 4

chromatography. The chiral auxialiary was then removed with NaOH for compounds 12a and 12b. Scheme 2 was carried out on each individual enantiomer to obtain each diastereoisomer and the same sequence of reactions described in acid to obtain diastereoisomers 11a and 11b which were separated by resolution. The alcohol 6 was coupled with a chiral Boc-protected amino Preparation of chiral analogs was performed through chemical

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the corresponding ketones from any of several methods known to those skilled in the art. Oximes of formula I or II wherein X is C=NOR4 are prepared from

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WO 00/66559 PCT/US00/11633

. 20 -

ಠ ഗ an acid under standard conditions to obtain a compound of formula I or II. aqueous HCl or trifluoroacetic acid, and the resulting amine is coupled to BOC protecting group is removed by treatment with an acid such as separated or the mixture carried through and separated at the end. The amine hydrochloride in the presence of a base such as sodium acetate. formula I and II, is dissolved in a solvent such as CH3OH or ethanol and The resulting mixture of Z- and E-O-substituted oximes 13 can be treated with an R4-substituted hydroxylamine such as O-methylhydroxyl-In Scheme 5, the ketone 1a, wherein R and R1 are as defined for

Scheme 6:

ᇙ oxime is then treated with a base such as potassium hexamethyldisilazide electrophiles, to yield the desired O-substituted oxime. agent, e.g., CH₃I, dimethylsulfate, CH₃CH₂I, trifluoroethyl triflate or similar in a suitable solvent such as DMF followed by treatment with an alkylating similar conditions to yield, after separation, the E- and Z-oximes. Each

Alternatively, the ketone 1s can be treated with HONH2·HCI under

8 known methods as shown in Schemes 7 and 8. The ketone starting material of formula 1a can be prepared by

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ð თ under standard conditions using BOC anhydride to give 1a. presence of a dehydrating agent such as titanium isopropoxide followed by ketal is removed by treatment with aqueous acid followed by re-protection CH3MgBr or vinylmagnesium bromide to give the alkylated product 22. The aminonitrile is treated with a grignard reagent (R1Mg-halide) such as treatment with diethylaluminum cyanide to give an aminonitrile 21. The resulting free amine 20 is treated with N-BOC-piperidine-4-one in the standard conditions. The N-trifluoroacetyl group is removed and the yields a ketone 18 which is converted to its ethylene ketal 19 under suitable catalyst such as AICl3 and optionally in a solvent such as CH2Cl2 isonipecotoyl chloride 17 and an aromatic group R-H in the presence of a In Scheme 7, Friedel-Crafts condensation of N-trifluoroacetyl-

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(Chen et al, <u>Tetrahedron Lett.</u>, <u>3</u>Z, 30 (1996), 5233-5234), is transformed to Alternatively, 23, prepared via Wittig olefination of N-BOC-piperidone

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WO 00/66559 PCT/US00/11633

TPAP/NMO to give the desired ketone. treated with an oxidizing agent such as Dess-Martin periodinane or suitable solvent such as ether or THF and the resulting alcohol 28 is aldehyde 27. The aldehyde is treated with an aryllithium reagent in a perruthenate (TPAP) and N-methylmorpholine N-oxide (NMO) to give with a suitable oxidant such as a mixture tetrapropylammonium converted to alcohol 26 by hydroboration/oxidation. Alcohol 26 is treated intermediate 25 by analogy to the procedure described in Scheme 7. 25 is

ಠ and R¹⁹ are the same, or wherein R and R¹⁹ are different are prepared procedures apply to other R and R19 groups. R is phenyl and R19 is CF3-phenyl, respectively, but the general exemplified by processes wherein R and R19 are each phenyl and wherein according to schemes 9 and 10, respectively. The schemes are Compounds of formula I or II wherein X is -C(R13)(R19)-, wherein R

Scheme 9

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않 8 procedure, e.g., treatment with a reagent R2COOH and coupling agents methylene bond is reduced using standard conditions to obtain the BOCwith TFA, and the resultant amine subjected to a standard amidation protected diphenylmethyl-piperidine of formula 46, the BOC group is obtain the BOC-protected diphenylmethylene-piperidine of formula 45. The compound of formula 44, which is then treated with phenylboronic acid to compounds 20-22 of Scheme 7, the BOC group is removed by treatment removed and the amine of formula 47 is treated as described for N-BOC-4-piperidone is treated with CBr4 to obtain the di-bromo

such as EDCI, HOBT and a base, to obtain the compounds of formula 48

WO 00/66559

PCT/US00/11633

- 23 -

N-BOC-4-piperidone is treated with a reagent such as diethyl benzylphosphonate to obtain the phenylmethylene-piperidine of formula 49, which is then brominated to obtain the bromophenylmethylene-piperidine of fomula 50. The BOC protecting group is removed using standard conditions, e.g., treatment with TFA, to obtain amine 51, and the amine 51 is treated as described for compounds 20-22 of Scheme 7 to obtain the aminonitrile 52, then the protected amine 53. The amine 53 is treated with a reagent such as 4-CF3-phenylboronic acid to obtain compound 54 and the methylene bond is reduced using standard conditions to obtain racemic 55. The BOC group is removed by treatment with TFA, and the resultant amine subjected to a standard amidation procedure, e.g., treatment with a reagent R²COOH and coupling agents such as EDCI, HOBT and a base, to obtain the racemic compounds of formula 56.

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Compounds useful in this invention are exemplified by the following preparative examples, which should not be construed to limit the scope of

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WO 00/66559 PCT/US00/11633

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the disclosure. Alternative mechanistic pathways and analogous structures within the scope of the invention may be apparent to those skilled in the art

5 A solution of free amine 29 (1.45 g, 3.97 mmol) and 2,6-dimethyl-benzoyl chloride (840 mg, 5.0 mmol) in aqueous 1 N NaOH (20 ml) and CH₂Cl₂ (20 ml) was stirred overnight at RT. The reaction mixture was extracted with CH₂Cl₂, dried over Na₂SO₄ and concentrated under high vacuum to provide 30 (1.97 g, 97%), as a slightly yellow foam.

10 To a solution of ketone 30 (550 mg, 1.11 mmol) in CH₃OH (6 ml) was added NaBH₄ (60 mg, 1.59 mmol) and the solution was stirred overnight at RT. The reaction mixture was then poured into 0.1 N NaOH extracted with CH₂Cl₂, dried over Na₂SO₄, and concentrated to give 31

15 Example 1A:

(543 mg, 98%), as a slightly yellow foam.

To a solution of alcohol 31 (50 mg, 0.10 mmol) in anhydrous DMF (0.5 ml) was added NaH (6.0 mg, 0.25 mmol) followed by ethyl iodide (12 µl, 0.15 mmol) and the reaction was stirred 4 h at 40 °C. The reaction mixture was poured into aqueous 0.1 N NaOH, extracted with CH₂Cl₂. 20 dried over Na₂SO₄, and concentrated. Purification by preparative chromatography (eluting with CH₂Cl₂/CH₃OH, 9:1) yielded 1A (31 mg, 59%) as a colorless oil: 1H ·NMR (300 MHz, CDCl₃) & 7.39 (br d, J = 8.4 Hz, 2H), 7.02-7.12 (m, 3H), 6.95 (m, 2H), 3.94 (m, 1H), 3.79 (d, J = 7.2 Hz, 1H), 3.10-3.35 (m, 4H), 2.60-3.00 (m, 3H), 2.19 (br s, 6H), 1.60-2.10 (m, 5H), 1.05-1.50 (m, 5H), 1.08 (br t, 3H), 0.94 (s, 3H); HRMS (MH+)

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Example 1B

(d, 3H), 1.60-2.10 (m, 5H), 1.05-1.45 (m, 5H), 1.08 (m, 3H), 0.95 (s, 3H); 7.05-7.15 (m, 3H), 6.97 (m, 2H), 5.40 (d, J = 7.8 Hz, 1H), 4.09 (m, 1H), a colorless oil: ^{1}H -NMR (300 MHz, CDCl₃) 5 7.42 (br d, J = 8.2 Hz, 2H), chromatography (eluting with CH₂Cl₂/CH₃OH, 9:1), 1B (44.7 mg, 81%) as 0.20 mmol) in anhydrous CH₂Cl₂ (0.5 mL) was added propionyl chloride 3.43 (m, 1H), 3.23 (m, 1H), 2.96 (m, 1H), 2.82 (m, 1H), 2.70 (m, 1H), 2.21 reaction mixture was treated as for 1A to give, after preparative (30 μ l, 0.30 mmol) and the solution was stirred overnight at RT. The To a solution of alcohol 31 (50 mg, 0.10 mmol) and pyridine (16.2 μ l

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ಠ Example 1C: To a solution of alcohol 31 (29.4 mg, 0.059 mmol) and HRMS (MH+) 555.2230.

8 5 (m, 1H), 2.82 (m, 1H), 2.70 (m, 1H), 2.22 (br s, 3H), 1.60-2.10 (m, 5H), 7.2 Hz, 1H), 4.09 (m, 1H), 3.71 (m, 3H), 3.45 (m, 1H), 3.24 (m, 1H), 2.97 Hz, 2H), 7.14 (d, J = 8.4 Hz, 2H), 7.09 (m, 1H), 6.98 (m, 2H), 5.21 (d, J =preparative chromatography (eluting with CH2Cl2/CH3OH, 9:1), 1C (15 mg overnight at RT. The reaction mixture was treated as for 1A to give, after pyridine (9.5 μ l, 0.118 mmol) in anhydrous CH₂Cl₂ (0.3 mL) was added 46%) as a colorless oil: 1 H -NMR (300 MHz, CDCl₃) δ 7.46 (br d, J = 8.4methylchloro-formate (13.8 μ l, 0.18 mmol) and the solution was stirred

(m, 5H), 0.95 (s, 3H); HRMS (MH+) 557.2017

Example 1D:

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2H), 2.20 (br s, 3H), 1.55-2.10 (m, 5H), 1.10-1.50 (m, 5H), 0.95 (s, 3H); d, J = 8.2 Hz, 2H), 7.05-7.15 (m, 3H), 6.98 (m, 2H), 5.34 (m, 1H), 4.08 (m, HRMS (MH+) 556.2169. 1H), 3.44 (m, 1H), 3.24 (m, 1H), 3.19 (s, 3H), 2.96 (m, 1H), 2.65-2.85 (m, give, after preparative chromatography (eluting with CH2Cl2/CH3OH, 9:1), was stirred 5 h at 45 °C. The reaction mixture was treated as for 1A to mmol) and methylisocyanate (40 μ l, 0.68 mmol) in anhydrous THF (0.3 m) 1**D** (25 mg, 75%) as a colorless oil: 1H -NMR (300 MHz, CDCl₃) δ 7.42 (br A solution of alcohol 31 (30 mg, 0.060 mmol), pyridine (9.7 μ l, 0.12

Example 1E:

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CH₂Cl₂/CH₃OH, 9:1), 1**E** (50 mg, 86%) as a colorless oil: ¹H -NMR (300 as for 1A to give, after preparative chromatography (eluting with (6 mg, 0.15 mmol), and 2-chloropyridine (28.2 μ l, 0.30 mmol) in anhydrous DMF (0.5 ml) was stirred 16 h at 90 °C. The reaction mixture was treated A solution of alcohol 31 (50 mg, 0.10 mmol), NaH 60% in mineral oil

> WO 00/66559 PCT/US00/11633

26-

3H); HRMS (MH+) 576.2230. d, J = 7.0 Hz, 1H), 4.09 (m, 1H), 3.44 (m, 1H), 3.24 (m, 1H), 2.65-3.05 (m 3H), 2.22 and 2.23 (s, 3H), 1.60-2.15 (m, 5H), 1.10-1.50 (m, 5H), 0.87 (s, 2H), 7.21 (d, J = 8.0 Hz, 2H), 6.95-7.15 (m, 3H), 6.65-6.80 (m, 2H), 5.74 (br MHz, CDCl₃) δ 7.98 (m, 1H), 7.47 (brt, J = 7.2 Hz, 1H), 7.38 (d, J = 8.0 Hz

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prepared Using similar procedures, compounds of the following structure were

| | | | | | | | | ō |
|-----------|---------------|----------|--|---|----------------------|---------------------------------------|---------------------|--|
| | ź | 1 | = | Ī | 16 | Ħ | , m | where |
| Br | 略 | 망 | Br | Br | Br | Вг | 공 | in R3, R6 |
| √ N N | بز \ \ الم | Z KN | -C(O)NHCH ₂ CH ₃ | -C(0)-(CH ₂) ₂ CH ₃ | -C(O)CH ₃ | -C(0)OCH ₂ CH ₃ | R3 | wherein R3, R6 and R2 are as defined in the table: |
| н₃с, фсн₃ | н₃с, Дсн₃ | ҸҙҀѼ҉СҸҙ | нзсДснз | н _з с Ёсн _з | нзсССнз | н _з с Снз | Ą | d in the table: |
| 577.2183 | 577.2162 | 584.1786 | 572.2322 | 569.2392 | 541.2054 | 571.2181 | HRMS (MH+) found | |

Additional data for compounds of Example

- 27 -

8.39 (d, J = 5.6 Hz, 1H), 7.42 (d, J = 8.4 Hz, 2H), 7.27 (d, J = 8.4 Hz, 2H), 7.05-7.20 (m, 2H), 6.99 (m, 2H), 6.84 (m, 1H), 5.70 (d, J = 7.8Hz, 1H), 4.11 (m, 1H), 3.43 (m, 1H), 3.25 (m, 1H), 2.65-3.05 (m, 3H), 2.23 and 2.25 (s, 3H), 1.55-2.10 (m, 5H), 1.10-1.50(m, 5H), 0.88 (br s, 3H)

A solution of ketone 32 (0.60 g, 1.29 mmol) and NaBH₄ (60 mg, 1.59 mmol) in CH₃OH (5 ml) was stirred overnight at RT. The reaction mixture was poured into 0.1 N NaOH, extracted with CH₂Cl₂, dried over Na₂SO₄, and concentrated to give 33 (0.60 g, 100%), as a white foam.

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To a solution of alcohol 33 (543 mg, 1.2 mmol) in anhydrous toluene (4 mi) was added KHMDA, 0.5 N in toluene (2.6 ml, 1.30 mmol) followed, 15 min. later, by 2-bromopyridine (125 μ I, 1.30 mmol). The reaction was heated 5 h at 60 °C, cooled to RT and poured into 5% aqueous NaHCO₃ (25 ml). Extraction with CH₂Cl₂, drying over Na₂SO₄ and concentration afforded an oil which was purified by flash chromatography over silica gel (eluting with CH₂Cl₂/AcOEVEt₃N 50:50:1 to 40:60:1) to yield 34a (310 mg,

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A solution of **34a** (310 mg, 0.57 mmol) in anhydrous CH₂Cl₂ (2 ml) and TFA (2 ml) was stirred 30 min. at RT. After concentration, the residue was taken up in aqueous 1 N NaOH, extracted with CH₂Cl₂, dried over Na₂SO₄ and concentrated to give **34b** (220 mg, 87%), as a white foam.

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49%), as a yellow foam.

WO 00/66559 PCT/US00/11633

A solution of free amine **34b** (85 mg, 0.19 mmol), 2,4-dimethylnicotinic acid (50 mg, 1.45 mmol), DEC (60 mg, 0.31 mmol), HOBT (50 mg, 0.37 mmol) and N-methylmorpholine (80 ml, 0.72 mmol) in anhydrous DMF (1 ml) was stirred overnight at 40 °C. After concentration, the residue was taken up in aqueous 0.1 N NaOH, extracted with CH₂Cl₂, and dried over Na₂SO₄. The residue obtained after concentration of the solvent was purified by preparative chromatography over silica gel (eluting with CH₂Cl₂/CH₃OH/NH₄OH, 96:4:1) to afford **35** (95 mg, 85%), as a colorless oil: 1H -NMR (300 MHz, CDCl₃) δ 8.33 (d, J = 5.1 Hz, 1H), 7.99 (dd, J = 4.8 and 1.8 Hz, 1H), 7.86 (d, J = 8.4 Hz, 2H), 7.56 (d, J = 8.4 Hz, 2H), 7.53 (m, 1H), 6.96 (d, J = 5.1 Hz, 1H), 6.75-6.85 (m, 2H), 4.15 (m, 1H), 3.45 (m, 1H), 3.02 (s, 3H), 2.99 (m, 2H), 2.79 (m, 1H), 2.47 and 2.48 (s, 3H), 2.45 (m, 1H), 2.25 and 2.26 (s, 3H), 1.65-2.15 (m, 5H), 1.15-1.55 (m, 5H), 0.90 (s, 3H); HRMS (MH+) 577.2858.

15 Using similar procedures, compounds of the following structure were prepared

| 2E | 2D | 2C | 28 | 2A | Ϋ́ | whe |
|------------------|------------------|--------------------------------|------------------|---------------------|------------|--|
| H3CSO2- | Br | Br | Br | B. | r. Re | rein H3, H3 |
| 2: N | Z | Z) | Z) | z) | P3 | and R2 ar |
| .γ. <u>Ο</u> | Н ₃ С | _Э | H ₃ C | ν _ζ ο | | e as define |
| ₩ _{NH2} | NH ₂ | C C N CH ₃ | Ç ₩ P | NH _N | R2 | wherein R3, R6 and R2 are as defined in the table: |
| 597.2296 | 577.2172 | 577.2172 | 578.2006 | 599.1062 | HRMS (MH+) | Φ. |

- 29 -

- 30 -

| 2EE | 2DD | 2CC | 288 | 2AA | 22 | 24 | 2X | 2W | 2V | 2U | 21 | 28 |
|----------|--|----------|----------------------------------|--|--------------------|---|----------------------------------|--|---------------------------------------|-------------------------------------|-----------------------------------|----------------------------------|
| Br | Ω | Ω | Ω | Ω | F ₃ CO- | F3CO- | F ₃ C. | H ₃ CSO ₂ - | H3CSO2- | H3CSO2- | H3CSO2- | H3CSO2- |
| \Z | \(\sigma_2\) | _=Z | , | | | \ | _{=\infty} | N. N. | Ş | N. Z | z | Z Z |
| CI | H ₃ C√√CH ₃ N≪N | CI CI | H ₃ C CH ₃ | H ₃ C CH ₃ N | CI | H ₃ C ₁ CH ₃ CH ₃ | H ₃ C CH ₃ | H ₃ C CH ₃ CH ₃ | H ₃ C N CH ₃ | H ₃ C CH ₃ | H ₃ C. CH ₃ | H ₃ C CH ₃ |
| 619.1062 | 534.2637 | 573.1606 | 549.2646 | 533.2673 | 623.1790 | 584.2848 | 583.2905 | 599.2362 | 576.2896 | 583.2426 | 594.2750 | 578.2801 |

| 2R | 20 | 2P | 20 | 20 | 2M | 2L | 25 | 22 | 22 | 27 | 26 | Ņ Ti |
|----------|------------------|----------|----------------------------------|----------|------------------------------------|--|----------|--------------------|---------------------------------------|-----------------------------------|----------------------------------|----------------------|
| H3CSO2- | Br | Ð | F ₃ CO- | Bŗ | Br | F3CO- | F3CO- | F ₃ CO- | H3CSO2- | H ₃ CSO ₂ - | F ₃ C ₋ | H3CSO ₂ . |
| , T, Z | χ _ν χ | z. | ų, N | Z Z | 4, | تر حر | | z | , , , , , , , , , , , , , , , , , , , | z | z) | Z |
| | | | H ₃ C CH ₃ | Н₃С₩ОН | H ₃ C → CH ₃ | H ₃ C N CH ₃ | н₃с₩он | н₃с Дсн₃ | H ₃ C CH ₃ | н₃с Сн₃ | H ₃ C CH ₃ | н₃СДон |
| 578.2792 | 594.2072 | 595.2114 | 599,2847 | 579.1986 | 580.2123 | 583.2913 | 584.2744 | 582.2969 | 593.2805 | 576.2890 | 567.2947 | 578.2697 |

31 -

| _ | | | | | | | | | | | | |
|---|--|------------------------------------|-----------|---------------------------------|---|---|----------------------------------|-------------------------------|---|-----------------------------------|---|---|
| | 200 | 2PP | 200 | 2NN | 2MM | 2LL | 2KK | 211 | 2 | 2HH | 2GG | 2FF |
| | Br | BF | π | ō | 71 | ті | т | F ₃ C ₂ | H3CSO2- | H ₃ CSO ₂ - | F ₃ C- | H3CSO2- |
| | \\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\ | H ₃ C ₂ N | 7 | 7 | , (z | | \ | \[\sum_{=\infty}^{\infty} \] | ~, Z= | | _Z_Z | S N |
| 2 | H ₃ C CH ₃ | H ₃ C CH ₃ | O+2 CΩ | 0 4 2 \\ \frac{1}{2} | H ₃ C ₁ CH ₃ | H ₃ C ₁ CH ₃ CH ₃ | H ₃ C CH ₃ | CI | H ₃ C ₁ CH ₃ | ZCI CI | H ₃ C ₁ CH ₃ | H ₃ C ₁ CH ₃ |
| | 607.2291 | 591.2330 | 573.1818 | 589.1534 | 518.2944 | 533.2916 | 517.2696 | 607.1871 | 579.2749 | 618.1722 | 568.2913 | 584.2375 |
| | | | | | | | | | | | | |

WO 00/66559 -32 -

PCT/US00/11633

| 2XX | 2WW | 2VV | 200 | 211 | 288 | 2RR |
|---|----------|-------------------|------------------------------------|-------------------|----------|---------------------------------------|
| I | 711 | F ₃ C- | F ₃ C- | F ₃ C. | Вг | Br |
| \\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\ | 14, Z | N. P. N. | \ | ₹ <u></u> | * | , Y |
| H ₃ C ₁ CH ₃ | CH3 CH3 | сн₃ Сн₃ | CH ₃ CH ₃ | 04Z } | oZ∑} | H ₃ C N CH ₃ |
| 500.3023 | 532.3106 | 567.2961 | 583.2909 | 623.1809 | 633.1040 | 592.2294 |
| | · | | | | | |

Additional data for compounds of Example 2:

| | The second secon |
|-----|--|
| Ēx. | 1H-NMR (300 MHz 1H NMR (CDCl3)) |
| 2A | 7.98 (m, 1H), 7.49 (br t, $J = 7.1$ Hz, 1H), 7.39 (d, $J = 8.1$ Hz, 2H), |
| | 7.12 (d, $J = 8.1$ Hz, 2H), 7.01 (t, $J = 8.4$ Hz, 1H), 6.65-6.80 (m, 3H), |
| - | 6.56 (d, $J = 8.4$ Hz, 1H), 5.76 (d, $J = 7.2$ Hz, 1H), 3.95-4.20 (m, 1H), |
| | 3.89 and 3.92 (s, 2H), 3.30-3.55 (m, 2H), 3.12 (m, 1H), 2.70-3.00 |
| | (m, 2H), 1.65-2.10 (m, 5H), 1.20-1.60 (m, 5H), 0.95 and 0.99 (s, 3H) |
| 26 | 8.31 (d, 1H), 8.01 (d, 1H), 7.50 (m, 4H), 6.95 (d, 1H), 6.80 (m, 2H), |
| | 5.90 (d, 1H), 4.15 (d, 1H), 3.25-3.55 (m, 2H), 2.80-3.15 (m, 3H), |
| | 2.50 (d, 3H), 2.30 (d, 3H), 1.80-2.15 (m, 7H), 1.20-1.60 (m, 5H), |
| | 0.92 (s, 3H) |

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2<u>8</u>2 2PP 2MM 2 1 1 29 2 쏫 | 8.17 (s, 1H), 8.01 (br d, J = 4.0 Hz, 1H), 7.50 (br t, J = 8.0 Hz, 1H)8.37 (d, J = 6.0 Hz, 1H), 7.83 (br d, J = 4.6 Hz, 1H), 7.41 (d, J = 8.4 Hz8.93 (s, 1H), 8.04 (br d, J = 4.8 Hz, 1H), 7.50 (m, 1H), 7.32 (m, 2H), 2.47 and 2.49 (s, 3H), 2.23 and 2.26 (s, 3H), 2.23 (s, 3H), 1.65-2.15 2H), 2.80 (m, 1H), 1.70-2.15 (m, 5H), 1.10-1.50 (m, 5H), 0.90 (s, = 6.8 Hz, 1H), 4.18 (m, 1H), 3.44 (m, 1H), 3.39 (m, 1H), 3.00 (m, 7.20-7.35 (m, 4H), 6.78 (t, J = 6.8 Hz, 1H), 6.71 (m, 1H), 5.80 (d, J8.49 (s, 2H), 8.26 (br s, 1H), 8.04 (br s, 1H), 7.80-7.95 (m, 3H), 7.53 (m, 5H), 1.15-1.55 (m, 5H), 0.90 (s, 3H) 1H), 4.20 (m, 1H), 3.20-3.50 (m, 2H), 2.97 (m, 2H), 2.78 (m, 1H) J = 4.6 Hz, 1H), 6.68 (br t, J = 6.0 Hz 1H), 5.89 (br d, J = 6.8 Hz, Hz, 2H), 7.34 (d, J = 6.0 Hz, 1H), 7.22 (d, J = 8.4 Hz, 2H), 6.97 (d, 3H), 1.90-2.15 (m, 3H), 1.70-1.90 (m, 2H), 1.15-1.50(m, 5H), 0.90(s 1H), 3.25-3.50 (m, 2H), 2.93 (m, 2H), 2.78(m, 1H), 2.44 and 2.46 (s, 6.97 (m, 2H), 6.78 (m, 1H), 6.72 (m, 1H), 5.82 (m, 1H), 4.21 (m, (m, 2H), 2.94 (m, 2H), 2.80 (m, 1H), 1.75-2.15 (m, 5H), 1.25-1.50 (d, J = 8.4 Hz, 2H), 5.81 (d, J = 6.8 Hz, 1H), 4.16 (m, 1H), 3.30-3.50 and 2.46 (s, 3H), 2.23 and 2.25 (s, 3H), 1.65-2.10 (m, 5H), 1.15-1H), 3.32 (m, 1H), 2.95 (m, 1H), 2.86 (m, 1H), 2.75 (m, 1H), 2.44 Hz, 2H), 7.25 (d, J = 8.0 Hz, 2H), 6.98 (d, J = 6.4 Hz, 1H), 6.78 (m, 8.14 (d, J = 6.8 Hz, 1H), 8.02 (m, 1H), 7.51 (m, 1H), 7.41 (d, J = 8.03H), 1.60-2.20 (m, 5H), 1.05-1.50 (m, 5H), 0.85 (br s, 3H) 3.26 (m, 1H), 2.65-3.05 (m, 3H), 2.41 and 2.42 (s, 3H), 2.20 (br s, 5.2 Hz, 1H), 5.67 (d, J = 7.2 Hz, 1H), 4.07 (m, 1H), 3.43 (m, 1H), 1H), 7.38 (d, J = 8.4 Hz, 2H), 7.18 (d, J = 8.4 Hz, 2H), 6.92 (d, J =8.29 (d, J = 5.2 Hz, 1H), 8.18 (m, 1H), 7.98 (br s, 1H), 7.89 (br s, 2.10 (m, 5H), 1.15-1.55 (m, 5H), 0.78 and 0.91 (s, 3H) 3.55 (m, 1H), 3.32 (m, 1H), 2.70-2.95 (m, 2H), 2.18 (s, 3H), 1.65-7.01 (m, 1H), 6.60-6.75 (m, 4H), 5.77 and 5.79 (d, J = 5.6 Hz, 1H) 7.97 (m, 1H), 7.45 (m, 1H), 7.32 (t, J = 8.4 Hz, 2H), 7.06 (m, 2 H).1H), 6.73 (m, 1H), 5.78 (d, J = 6.8 Hz, 1H), 4.17 (m, 1H), 3.43 (m, 1.50 (m, 5H), 0.90 (s, 3H) 5H), 0.89 (s, 3H)

> WO 00/66559 PCT/US00/11633

-34

5 ഗ and Z isomers. and chromatographed to yield 1.50 g (94%) of oxime 36, as a mixture of E extracted with CH2Cl2. The combined extracts were dried, concentrated 24 h. The resulting mixture was then poured into aqueous NaOH and hydrochloride (3.26 g, 47 mmol), and the solution was stirred at RT for added sodium acetate (5.0 g, 47 mmol) and C-Methyl hydroxylamine To a solution of ketone 30 (1.5 g, 3.22 mmol) in CH₃OH (50 ml) was

ᇙ EtOH/EtOAc) to afford 0.100 g (50%) of amine 37. concentrated and chromatographed over silica gel (eluting with 20% water and extracted with CH₂Cl₂. Combined organic layers were reaction was warmed slowly to 60°C for 2 h, cooled to RT, quenched with was added BH3•THF (1.0 M solution in THF) at 0 °C and the solution was cooled to 0°C and a solution of 1N KOH in CH3OH (5 ml) was added. The then warmed to RT and stirred for 1 h. The reaction mixture was then To a stirred solution of oxime 36 (0.200 g, 0.380 mmol) in THF (5 ml)

8 desired product 38: 1H-NMR (300 MHz, CDCl₃) § 7.45 (d, 2H), 7.05-7.12 concentrated and purified by preparative chromatography to give 0.010 g of overnight. It was then poured into water, extracted with EtOAc, dried, pyridine (0.5 ml) and CICOOCH₃ (0.25 ml), and the solution was stirred (m, 3H), 6.95 (d, 2H), 4.95 (m, 1H), 4.45 (m, 1H), 4.15 (m, 1H), 3.62 (s, To a stirred solution of amine 37 (0.015 g, 0.030 mmol) was added

(m, 12H), 0.90 (s, 3H); HRMS (MH+) 558.3013. 3H), 3.47 (m, 1H), 3.25 (m, 1H), 2.88-3.10 (m, 3H), 2.25 (s, 6H), 1.20-2.10

Example 4

Enantiomer

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concentration of the solvent was subjected to flash chromatography over silica gel (eluting with CH₂Cl₂/acetone, 9:1) to afford, in order of elution: (i) first 40a (391 mg, 38%), as a white foam; (ii) second 40b (391 mg, 38%), with CH₂Cl₂, and dried over Na₂SO₄. The residue obtained after reaction mixture was poured into aqueous saturated NaHCO3, extracted mmol) in anhydrous CH2Cl2 (5 ml) was stirred overnight at RT. The (413 mg, 1.50 mmol), DEC (290 mg, 1.50 mmol) and DMAP (190 mg, 1.55 A solution of alcohol 39ab (660 mg, 1.41 mmol), Boc-Thr(t-Bu)-OH

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as a white foam.

20 ភ 43b (Enantiomer B.)). gave 39b (Enantiomer B). 40a gives 43a (Enantiomer A) and 40b gives A) (246 mg, 98%) as a white foam. (Following the same procedure, 40b aqueous 0.1 N NaOH and extracted with CH2Cl2 to yield 39a (Enantiomer was stirred at 65 °C for 3 h. The final mixture was then poured into (3 ml) was added NaOH (110 mg, 2.75 mmol; 5 equiv.) and the solution To a solution of diastereoisomer 40a (391 mg, 0.54 mmol) in CH₃OH

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WO 00/66559

PCT/US00/11633

poured into aqueous sat'd NaHCO3, extracted with CH2Cl2, dried over anhydrous DMF (1.5 ml) was stirred 2 h at 75 °C. The reaction mixture was oil (23 mg, 0.96 mmol), and 2-bromopyridine (60 μ l; 0.62 mmol) in A solution of alcohol 39a (210 mg, 0.45 mmol), NaH 60% in mineral

CH2Cl2/AcOEt/Et3N, 60:40:0.5 to 40:60:0.5) to afford 41a (143 mg, 59%) Na₂SO₄ and purified by flash chromatography over silica gel (eluting with Removal of the Boc-protecting group in 41a (93 mg, 0.17 mmol)

proceeded as for 34b to provide 42a (68 mg, 91%), as a white foam. The amine 42a (50 mg, 0.11 mmol) was coupled with 4,6-dimethyl-

ᇙ ö 3H)); HRMS (MH+) 578.2140. 8.92 (s, 1H), 8.02 (m, 1H), 7.51 (m, 1H), 7.51 (brt, J = 8.4 Hz, 1H), 7.41 (d pyrimidine-5-carboxylic acid following the conditions described for the synthesis of 35 to yield 43a (28 mg, 44%). 1H-NMR (300 MHz, CDCi3) & 1H), 2.44 and 2.46 (s, 3H), 1.65-2.15 (m, 5H), 1.15-1.50 (m, 5H), 0.90 (s, (m, 1H), 4.19 (m, 1H), 3.41 (m, 1H), 3.36 (m, 1H), 2.94 (m, 1H), 2.78 (m, J = 8.4 Hz, 2H), 7.24 (d, J = 8.4 Hz, 2H), 6.78 (m, 1H), 6.73 (m, 1H), 5.78

The following compounds were prepared via similar methods:

wherein R3 R6 and R2

| Wilele | יייי חיי | AIIO The BIE | wilelelli no, no allo no ale as dell'ied il tile table. | the lable. | |
|---------|-----------|--------------------|---|---|---------------------|
| , m | Enantiome | R ⁶ | R3 | R² | HRMS (MH+) found |
| 4A | Α | Br | N. N. | н₃С Д сн₃ | 577.2172 |
| 4B | В | Br | N | н₃с Д сн₃ | 577.2162 |
| 40 0 | В | Br | N | H ₃ C ₁ CH ₃ | 578.2119 |
| 4D | A | F ₃ CO- | Z, | H ₃ C ₁ CH ₃ | 584.2864 |

£

σ

583.2904

583.2862

- 37 -

| | 6 | \$ | \$ | 4 | \$ | 4 | 4 | ‡ | 46 | |
|--|--------------------|---|--|---------------------------------|--|---------------------------------|--------------------|--|---------------------------------------|--------|
| | œ | > | В | ω | Þ | > | 8 | Þ | . > | |
| | F ₃ CO- | F3CO- | Ω | Ω | Ō | Ω | F ₃ CO- | F ₃ CO- | F ₃ CO- | |
| حر در/ | \supset | z.) | \\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\ | _Z | \\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\ | ų/=z | , S | 1, | , S | 2, |
| N N N N N CH ₃ | -* | H ₃ C N N NH ₂ NH ₂ | H ₃ C N CH ₃ CH ₃ | CH ₃ CH ₃ | H ₃ C N CH ₃ | CH ₃ CH ₃ | H3C CH3 | H ₃ C N CH ₃ | CH ₃ ○+√CH ₃ | ر ح |
| | 599.2947 | 599.2947 | 548.2784 | 534.2644 | 548.2784 | 534.2639 | 598.3000 | 598.2994 | 599.2857 | |
| | | | | <u></u> | L | | | | | |

WO 00/66559 PCT/US00/11633

- 38 -

Additional data for compounds of Example 4:

| Ex. | ¹ H-NMR (300 MHz ¹ H NMR (CDCl ₃)) |
|-----|--|
| 46 | 8.05 (m, 1H), 7.97 (s, 2H), 7.53 (t, $J = 7.5$ Hz, 1H), 7.41 (d, $J = 8.4$ |
| | Hz, 2H), 7.16 (d, $J = 8.4$ Hz, 2H), 6.81 (t, $J = 6.4$ Hz, 1H), 6.76 (m, |
| | 1H), 5.87 (m, 1H), 4.19 (m, 1H), 3.30-3.50 (m, 2H), 2.99 (m, 2H), |
| | 2.79 (m, 1H), 2.20 and 2.22 (s, 3H), 1.70-2.15 (m, 5H), 1.15-1.50 |
| | (m, 5H), 0.91 (s, 3H) |
| 4 | 8.03 (m, 1H), 7.53 (m, 1H), 7.39 (d, $J = 8.4$ Hz, 2H), 7.14 (d, $J = 8.4$ |
| | Hz, 2H), 6.79 (t, $J = 6.8$ Hz, 1H), 6.73 (m, 1H), 5.87 (m, 1H), 4.19 |
| | (m, 1H), 3.42 (m, 1H), 3.37 (m, 1H), 2.98 (m, 2H), 2.80 (m, 1H), |
| | 2.41 and 2.43 (s, 3H), 1.90-2.15 (m, 3H), 1.70-1.90 (m, 2H), 1.20- |
| | 1.50 (m, 5H), 0.91 (s, 3H) |

and

- თ Excess TFAA is removed under vacuo, the reaction mixture is taken up in g of this amide is treated with SOCl₂ (300 ml) and the reaction mixture EtOAc, washed with water and concentrated to give 160 g of the amide. 50 acid (96 g) at 0°C and the reaction mixture is heated at reflux for 4h. Trilluroacetic anhydride (TFAA) (300 ml) is added to isonipecotic
- ಠ heated at reflux overnight. Excess thionyl chloride is then removed under vacuo to give 54 g of the acid chloride.
- 햐 g) in bromobenzene (40 ml) at ambient temperature and the reaction mixture is heated at reflux for 4 h. It is then cooled and poured into a AICl₃ (11g) is added slowly to a solution of the product of step 1 (10
- solution and concentrated to give 16.21 g of the desired ketone. mixture of conc. HCl and ice, and the product is extracted with EtOAc. The organic layer is separated and washed with water, half saturated NaHCO3 The product of step 2 (16.21 g) is dissolved in toluene (200 ml)
- 8 g of the desired ketal. further water is collected. The reaction mixture is concentrated to give 17.4 reaction mixture is heated at reflux with azeotropic removal of water until no containing ethylene glycol (25 ml) and p-toluenesulfonic acid (0.5 g). The

- and to this is added water (25 ml) and K2CO3 (12 g) and the reaction diluted with water and extracted with EtOAc. The organic layer is mixture is stirred at ambient temperature overnight. The reaction mixture is The crude product of step 3 (17.4 g) is dissolved in CH₃OH (100ml)
- of the desired amine. separated, washed with water and brine, and concentrated to give 12.55 g

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- BOC-piperidine-4-one (4.8 g, 24 mmol) in 1,2-dichloroethane (20 ml) is added titanium isopropoxide (6.7 ml, 32.3 mmol) and the mixture is stirred To a stirred solution of the product of step 4 (7.2 g, 23 mmol) and N-
- ಠ afford 7.3 g (63%) of the desired cyanide. for 12 h at RT. The reaction mixture is concentrated and a 1.0 M solution stirred for 2 h. The mixture is then filtered through celite and the resulting reaction mixture is then diluted with EtOAc, quenched with water (5 ml) and of diethyl aluminium cyanide (35 ml) is added at RT and stirred for 3 h. The filtrate is concentrated and chromatographed with 30 % EtOAc/hexanes to
- quenched with saturated aqueous NH₄Cl and extracted with CH₂Cl₂. The mmol) at RT and the mixture is stirred for 2 h. The reaction mixture is then THF (100 ml) is added a 3.0M solution CH_3MgBr in Et_2O (14.0 ml, 42 To a stirred solution of the product of step 5 (7.3 g, 14.03 mmol) in

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with EtOAc, dried and concentrated to yield 5.0 g (98%) of amine. h. The reaction mixture is then neutralised with 20%NaOH and extracted (40 ml) and conc. HCl (10 ml) is added and the mixture stirred at RT for 24 extracts are concentrated to afford 7.0 g of desired methylated compound The crude ketal of step 6 is dissolved in EtOAc (100 ml) and 6 N HC

8

- 25 EtOAc/hexanes to yield 5.1 g (79%) of the desired product. washed with brine, dried, concentrated and chromatographed with 20% stirred at RT overnight. The layers are separated and the organic layer is Et₂O (200 ml) is added 10% NaOH (50 ml) and BOC₂O, and the mixture is To a stirred solution of the product of step 7 (5.0 g, 13.6 mmol) in
- မ resulting mixture is then poured into aqueous NaOH and extracted with hydroxylamine hydrochloride and the mixture is stirred at RT for 24 h. The CH₃OH (50 mt) is added sodium acetate (5.0 g, 47 mmol) and O-Methyl CH₂Cl₂. The combined extracts are dried, concentrated and To a stirred solution of the product of step 8 (1.5 g, 3.22 mmol) in
- chromatographed to yield 1.5 g (94%) of oxime as a mixture of E and Z

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CH₂Cl₂ (10 ml) is added TFA (3 mL) and the mixture is stirred at RT for 2 h. To a stirred solution of the product of step 9 (1.5 g, 3.0 mmol) in

> WO 00/66559 PCT/US00/11633

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afford 1.2 g (100%) of amine. extracted with CH2Cl2. The combined extracts are dried concentrated to The reaction mixture is concentrated and poured into 10% NaOH and

CH₂Cl₂ is added 2,6-dimethylbenzoic acid (0.74 g, 4.96 mmol), EDCI (0.94 and concentrated to yield 1.6 g of oxime as a mixture of E and Z isomers. and the mixture is stirred for 12 h at RT. The reaction mixture is quenched g, 4.94 mmol), DIPEA (0.84 g, 6.58 mmol) and HOBT (0.66g, 4.94 mmol) with NaHCO3 and extracted with CH2Cl2. The combined extracts are dried To stirred solution of the product of step 10 (1.3 g, 3.2 mmol) in

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The isomers are separated by chromatography by eluting with CH2Cl2: Et2C

- ᇙ 3.00 (m, 3H), 2.82 (m, 1H), 2.24 (s, 3H), 2.23 (s, 3H), 2.15 (m, 3H), 1.80-E isomer: 300 MHz-1H NMR (CDCl3) & 7.5 (d, 2H), 7.23 (m, 2H), 7.10 (m 1.20 (m, 5H), 0.92 (s, 3H); MS FAB+ observed= 526.2070; estimated = (4:1) to afford 0.77 g of E isomer and 0.49 g of Z isomer. 1H), 6.90 (d, 2H), 4.03 (m,1H), 3.90 (s, 3H), 3.55 (m, 1H), 3.20 (m, 3H),
- 8 Z isomer: 300 Mhz - H NMR (CDCl₃) § 7.50 (d, 2H), 7.15 -6.95 (m, 5H), 3H), 2.25 (s, 3H), 2.10 (m, 2H), 1.80- 1.50 (m, 7H), 0.92 (s, 3H); 4.15 (m, 1H), 3.80 (s, 3H), 3.45 (s, 3), 3.25 (s, 3H), 3.00 (m, 2H), 2.24 (s,
- MS FAB+ observed= 526.2072; estimated = 526.2069

The following compounds were prepared via similar methods:

wherein X. R6 and R2 are as defined in the table:

| Wildigit A, D | מועק | שוופופווו A, חד מווע חד מופ מט עפוווופע ווו וויפ ומטופ. | נוזק ומטוק. | |
|---------------------|------|---|--|---------------------|
| χ | R | × | R ² | HRMS (MH+) found |
| 5A (mixture E/Z) | Br | —с— Ч-осн ₃ | H ₃ C\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\ | 529.1017 |
| 5B (mixture E/Z) | 8r | N,OCH3 | CI NH ₂ | 549.1023 |
| 5C | Br | сн _з сн ₂ о, N —с— | нзсфонз | 542.2210 |
| 5D | Br | —Ç— N-0CH3 | CI NH2 | 549.1011 |

- 41 -

- 42 -

| | Z | - - | | |
|----------------------|-------------------------|--|----|-----|
| Br L | , Ž | H ₃ CO N | ᄧ | 5EE |
| , J ^{Br} | | CH3CH2O N —C— | Br | 5DD |
| CH ₃ | н₃сү | H3CO_N —C— | Br | 5CC |
| CH ₃ | н₃Сү | H ₃ C N | 뫈 | 5BB |
| CH ₃ | H ₃ C√ N. | -c- - N | Вг | 5AA |
| N Br | Br | СН ₃ СН ₂ О, N —С— | Вг | 52 |
| CH ₃ | НзС | H ₃ C O | Br | 5Υ |
| 9-€ | - 2 H | сн ₃ сн ₂ о, -с- | Br | 5X |
| CH ₃ | НзС | CH3CH2Q C- | Br | 5W |
| KH3 CH3 | Н3С | CH ₃ (CH ₂) ₂ O, N —C— | Br | 5V |
| CH ₃ | Нзс | CH ₃ (CH ₂) ₂ Q N —C— | Br | 5U |
| CI N | 20 | H3CO_N | Br | 51 |
| 0+2/} | £ | CH3CH2O N | Br | SS |

| | 5R | 50 | 5P | 50 | 52 | 5M | 5L | 5 X | 5 | 51 | 5H | 5G | 5F | SE |
|---|-----------------------|-----------------------------------|---|---|--|----------------------------------|--|------------------|---|---|---|----------------------------------|----------|--|
| | 쯗 | Ē | ᄧ | 면 | ᄧ | Βŗ | Вг | Br | ₽r | B | Br | B | Ē | Br |
| | CH3CH2Q N —C— | CF3CH2O N —C— | CF3CH2Q N -C- | CH ₃ CH ₂ Q N -C- | CH3CH2Q N C | -с Р Осњенз | H ₃ C0 | CH3CH2Q N | H ₃ CO, | CH ₃ CH ₂ O, | сн ₃ сн ₂ о N —с— | H ₃ CO/N | H3CO_N | N OCH3 |
| 2 | н _э с-Хон, | H ₃ C√√CH ₃ | H ₃ C _N CH ₃ | CC CC | H ₃ C√√CH ₃ N≪N | H ₃ С√СН ₃ | H ₃ C√√CH ₃ N≪N | H ₃ C | H ₃ C _N CH ₃ | H ₃ C _N CH ₃ | н₃с√он | H ₃ C NH ₂ | н₃с√он | H ₃ C\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\ |
| | 541.2188 | 596.1831 | 595.1895 | 583.1061 | 542.2132 | 541.2194 | 528.1971 | 543.1000 | 527.2787 | 541.2178 | 542.1997 | 529,1017 | 530.1020 | 529.1128 |

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| - | | | | | 1 | | | | 1 | L | | | | | | 1 | 1 | | |
|---|---------------|-----------------------------------|----------|-----------------------------------|----------|----------|-------------------|-------------|---------------------|---------|-----------------------------------|----------|---------------------|----------|------------------|--------------|----------------|----------------------------------|-----------------------------------|
| | 600.2788 | | 616.2746 | | 610.2016 | | 555.2341 | 567.2327 | 555.2336 | | 517.2812 | 547.2902 | | 531.2956 | 546.3056 | | 556.2290 | | 686.9989 |
| | | | | | | | | | | | | | | | | | | | |
| | | | | | | | | | | | | | | | | | | | |
| | | | | | | | | | | | | | | | | | | | |
| | | 5AC | | 5AB | | 522 | | 544 | 5XX | | 5WW | | 5٧٧ | 500 | | 511 | | 5SS | 5RR |
| | | ᄪ | | Br | | Br | | Ψ | Ξ, | | Вг | | 막 | Φ | | Βr | | ᄪ | В |
| |) ()* | CH ₃ CH ₂ O | - Ci | CH ₃ CH ₂ O | Č | H3CO N | 0=z | сн₃сн₂о | C=Z | CH-CH-C | CH ₃ CH ₂ O | ೧=2 | н ₃ СО N | | 0 1 0 1 | CH3CH2O N | | снзсн20 | CH ₃ CH ₂ O |
| | o ch O CH3 | н₃СДХСН₃ | z)-{ | нэс-ү-снэ | z) | нэс-Жснз | V CH ₃ | _ {} | HyC CH ₃ | NH NH | H3C√. | 요~~~ | #. }-#. ⊊ | } | | ~2°H | SC SC SC | H ₃ C CH ₃ | N.O CH3 |
| | | 640.2868 | | 617.2488 | | 603.2349 | | 566.2407 | | 655 287 | 555.2336 | | 542.2002 | | 550 210 | 627.1729 | | 590.1995 | 593.2131 |

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- 43 -

WO 00/66559 PCT/US00/11633

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Additional data for compounds of Example 5:

ഗ A) Preparation of intermediate 27 (Scheme 8 (R1 = CH₃))

concentrated aqueous HCI (80 ml) for 1.5 h. The solution is concentrated, separated and the organic layer is extracted once with H₂O (20 ml). diluted with Et₂O (300 ml) and H₂O (150 ml), the aqueous layer is 23 (40.0 g, 0.203 mol) is vigorously stirred in EtOAc (200 ml) and

5 Combined aqueous layers are concentrated and the residue is dried 24 h under high vaccum to provide 26.7 g (84%) of a white solid. To this anhydrous CICH2CH2Ci (80 mL) with 4 Å molecular sieves, are hydrochloride and N-tert-butoxycarbonyl-4-piperidone (43.8 g, 0.22 mol) in

8 ಭ Celite (50 g). The reaction mixture is cooled to 0 °C, water (40 ml) is added the excess of water is quenched with Na₂SO₄. The final mixture is then slowly with vigorous stirring and, after an additional 5 min. stirring at RT, and is stirred overnight at RT. The mixture is then cooled to 0 °C and diethylaluminum cyanide, 1 N in toluene (260 ml, 0.26 mol) is added with (65.5 ml, 0.22 mol) at 0° C, the reaction mixture is allowed to warm to RT successively added DBU (33.2 ml, 0.22 mol) and titanium(IV) isopropoxide additional $3\, h$, after which are added CH $_2$ Cl $_2$ ($300\, m$ I), EtOAc ($300\, m$ I), and vigorous stirring. The reaction is allowed to warm to RT and stirred an

> WO 00/66559 PCT/US00/11633

as a colorless oil which solidifies upon standing. silica gel (eluting with Hexanes/EtOAc, 8:2), to provide 50.3 g (83%) of 24 filtered over Celite, evaporated and subjected to flash chromatography over

- ഗ at 0 °C is slowly added CH3MgBr 3 M in Et2O (91 ml, 3 equiv.) with vigorous stirring. After the addition, the reaction is allowed to warm to RT To a solution of 24 (27.7 g, 90.6 mmol) in anhydrous THF (200 mL)
- and stirred 3 h. The reaction is then poured into aqueous saturated NH₄C concentrated to give 27.1 g (100%) of 25 as a colorless oil. extracted with Et₂O (4 times), washed with brine, dried over Na₂SO₄, and
- ಠ 0 °C is slowly added BH3·S(CH3)2 2 N in THF (14 ml, 28 mmol) and the at 0 °C, 50 ml of a pH 7 buffer solution are added, followed slowly by 30% ml and slowly poured into ice-cooled EtOH/THF 1:1 (50 ml). After 15 min. H₂O₂ aqueous solution (50 ml). The reaction mixture is stirred overnight at solution is stirred 2 days at RT. The final mixture is concentrated to ca. 50 To a solution of 25 (11.6 g, 39.3 mmol) in anhydrous THF (50 ml) at
- ᇊ chromatography over silica gel (eluting with EtOAc/EtOH, 8:2) to yield 9.69 g (79%) of 26 as a colorless oil layers are dried over Na₂SO₄, concentrated, then subjected to flash RT, diluted with 1 N NaOH and extracted with CH₂Cl₂. Combined organic
- 25 8 oxide (4.67 g, 39.4 mmol) in anhydrous CH₂Cl₂ (100 ml) is stirred 1 h at is allowed to warm to RT and stirred 1 h. Additional N-methyl-morpholine A solution of 26 (11.2 g, 35.8 mmol) and N-methylmorpholine N-
- (eluting with CH₂Cl₂/acetone, 8:2 to 7:3) to provide 5.91 g (53%) of **27** as a concentrated, then subjected to flash chromatography over silica gel reaction to completion after 1 h. The reaction mixture is filtered over Celite, RT, cooled to 0 °C, and TPAP (885 mg) is added portionwise. The reaction N-oxide (1.30 g, 11 mmol) and TPAP (300 mg) are then added to drive the
- B) Preparation of title compounds of Example 6.
- မ aldehyde 27 (6.20 g, 20.0 mmol) in anhydrous THF (15 ml) is added allowed to warm to -50 °C for 10 min, cooled to -78 °C, and a solution of hexanes (11.2 ml, 28.0 mmol) is added via syringe. The reaction mixture is mmol) in anhydrous THF (100 mL) is cooled to -78 °C and n-BuLi 2.5 N in A solution of 1-bromo-4-(trifluoromethoxy)-benzene (4.20 ml, 28.0
- မ္ဟ dropwise. After stirring 30 min at -78 °C, then 30 min at -20 °C, the solution (94%) of an alcohol as a yellow oil. organic layers are dried over Na₂SO₄, and concentrated to give 8.85 g is poured into half-brine and extracted with CH₂Cl₂ (3 x 100 ml). Combined

WO 00/66559 - 47 - PCT/US00/11633

2) To a solution of the product of step 1 (8.85 g, 39.3 mmol) in CH₂Cl₂ (100 ml) at 0 °C is added Dess-Martin periodinane (19.70 g, 2.5 equiv.) and the reaction mixture is stirred 2 h at RT. An additional 8.0 g of Dess-Martin periodinane is added and the reaction is stirred for an additional

4 h. The solution is poured into a 1:1 mixture of aqueous saturated NaHCO₃ and aqueous saturated Na₂S₂O₃ (200 ml), stirred 10 min, extracted with CH₂Cl₂, and dried over Na₂SO₄. The residue obtained after concentration of the solvents is purified by flash chromatography over silica gel (eluting with hexanes/EtOAc, 7:3) to yield 5.48 g (63%) of the ketone as a yellow oil.

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3) A solution of the product of step 2 (2.85 g, 6.05 mmol), HONH₂·HCl (2.08 g, 30 mmol), and AcONa (2.46 g, 30 mmol) in EtOH (50 mL) is heated at reflux under N₂ for 4 h. After evaporation of the solvent, the residue is taken up in aqueous 0.1 N NaOH and extracted with CH₂Cl₂.

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- The residue obtained after evaporation of the solvents is subjected to flash chromatography over silica gel, to afford first the E-hydroxime (eluting with CH₂Cl₂/EtOAc, 7:3; 0.84 g; 29%), then the Z-hydroxime (eluting with CH₂Cl₂/EtOAc 1:1; 1.10 g; 37%), both products as white solids.
- 4) To a suspension of Z-hydroxime (0.89 g, 1.84 mmol) in anhydrous 20 DMF (5 ml) is slowly added KHMDA 0.5 N in toluene (4.0 ml, 2.02 mmol) at 0 °C, leading to the appearance of a yellow solution. After 2 min. at this temperature, dimethylsulfate (350 μl, 3.7 mmol) is slowly added and the solution is allowed to warm to RT and stirred 1 h. The mixture is poured into aqueous 0.1 N NaOH, extracted with CH₂Cl₂, and dried over Na₂SO₄.
- 25 The residue obtained after concentration of the solvents is purified by flash chromatography over silica gel (eluting with hexanes/EtOAc, 75:25) to afford 0.55 g (62%) of the Z-methoxime as a slighly yellow oil.
- A solution of Z-methoxime (0.59 g, 1.18 mmol) in anhydrous CH₂Cl₂ (6 ml) and TFA (3 ml) is stirred 1 h at RT. After concentration, the residue
 is taken up in aqueous 1 N NaOH, extracted with CH₂Cl₂, dried over
 Na₂SO₄ and concentrated to give 0.47 g (100%) of the free amine as a white foam.
- 6) A solution of the product of step 5 (470 mg, 1.18 mmol), 2,4-dimethylnicotinic acid (220 mg, 1.45 mmol), DEC (280 mg, 1.45 mmol), HOBT (243 mg, 1.80 mmol) and N-methylmorpholine (0.33 ml, 3.0 mmol) in anhydrous DMF is stirred 14 h. After concentration, the residue is taken up in aqueous 0.1 N NaOH, extracted with CH₂Cl₂, and dried over Na₂SO₄.

The residue obtained after concentration of the solvent is purified by flash

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WO 00/66559 PCT/US00/11633

- 48 -

chromatography over silica get (eluting with CH₂Cl₂/acetone, 7:3 to 1:1) to afford 640 mg (100%) of a colorless oil.

1H-NMR (400 MHz, CDCl₃) δ 8.35 (d, J = 7.8 Hz, 1H), 7.25 (AB system, 4H), 6.98 (d, J = 7.8 Hz, 1H), 4.22 (m, 1H), 3.82 (s, 3H), 3.43 (m, 1H), 3.33 (m, 1H), 2.99 (m, 2H), 2.85 (m, 1H), 2.49 (s, 3H, atropisomer a) and 2.51

HRMS (M+H+) 533.2747.

Following steps B-4, B-5, and B-6 using the E-oxime yields the

(s,3H, atropisomer b), 2.26 (s, 3H, atropisomer a) and 2/28 (s, 3H, atropisomer b), 1.95-2.21 (m, 3H), 1.20-1.90 (m, 7H), 0.92 (s, 3H),

The following compounds are prepared via similar procedures:

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corresponding E-methoxime product.

erein R4 R6 and R2 are as defined in the table

| wherein R4 | | R ⁶ and R ² are as defined in the table: | ed in the table: | |
|------------|--------------------|--|------------------|---------------------|
| Ř | | 72 | R ² | HRMS (MH+) found |
| 6A | 롸 | H ₃ C \{ | н₃С√СН₃ | 554.3000 |
| 68 | Br | H ₃ C∕—{ | нус Доснь | 555.2335 |
| గ్ర | Br | H ₃ C - { | н₃СДОН | 556.2175 |
| 69 | Br | насо Эм | HSC CH5 | 571.2284 |
| 39 | Br | н _э со | н₃С√Сн₃ | 570.2331 |
| କୁ | Br | <u></u> | нос Стонь | 569.1000 |
| 99 | F3CO- | | 40 Y CH | 601.2628 |
| Н9 | F ₃ CO- | | For F | 617.2549 |
| | | *** | ć | |

| 67 | ల | 67 | 6S | 6 P | ది | ę | 60 | 62 | M9 | £ | 65 | ผ | 61 |
|-------------------|---|------------|---|-----------------------------------|-----------|-----------------------------------|-------------------|--|---|--------------------|---|--------------------|---|
| F ₃ C- | F ₃ C- | Ω | F ₃ C- | Ω | Ω | Ω | F ₃ C- | Ω | Ω | F ₃ CO- | F ₃ CO- | F ₃ CO- | F3CO- |
| снзсн2- | СН ₃ - | снзсн2- | СН3- | CH ₃ CH ₂ - | CH3CH2- | CH ₃ CH ₂ - | СН3- | СН3- | сн _з сн ₂ - | کری | <u>ب</u> | الم | -Сн ₃ |
| Z | CI-\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\ | CI N CI | H ₃ C _H CH ₃ | H ₃ C CH ₃ | HC (4, CH | H ₃ C CH ₃ | HC A CH | H ₃ C N CH ₃ | H ₃ C _N CH ₃ | \$ G | H ₃ C N N N N N | н₃сДон | H ₃ C ₁ CH ₃ CH ₃ |
| 571.1838 | 557.1680 | 537.1603 | 518.2749 | 498.2633 | 513.2642 | 497.2683 | 533.2758 | 483.2516 | 513.2633 | 589.3013 | 602.2579 | 602.2465 | 534.2708 |

WO 00/66559 PCT/US00/11633

-50-

| | | | · · · · · · · · · · · · · · · · · · · | | | |
|---|-----------------|-----------------------------------|---|---|-----------------------------------|-----------------------------------|
| Addition | | 6AA | 62 | 67 | 6X | 6W |
| nal data fo | | F ₃ CO- | F ₃ CO- | F ₃ CO- | CI | CI |
| Additional data for compounds of Example 6: | | CH3CH2- | сн _з сн ₂ - | СН,- | CH ₃ CH ₂ - | CH ₃ CH ₂ - |
| kample 6: | NH ₂ | н ₃ СДХСН ₃ | H ₃ C ₁ CH ₃ N N N CH ₃ | H ₃ C N N OH ₃ | нас ДСНа | 0-2 C |
| | | 563.2939 | 562.3017 | 548.2853 | 497.2682 | 555.8401 |
| | | | | | | |

| × | Ex. 11H-NMR (300 MHz 1H NMR (CDC(3)) |
|----|--|
| 유 | 8.31 (d, 1H), 7.51 (d, 2H), 7.10 (d, 2H), 6.95 (d, 2H), 4.20 (m, 2H), |
| | 3.40 (d, 2H), 3.30 (m, 2H), 3.35 (m, 3H), 2.80-3.05 (m, 5H), 2.45 (d, |
| | 3H), 2.25 (d, 3H), 1.25-2.20 (m, 10H), 0.50 (m, 2 H), 0.22 (m, 2H), |
| | 0.90 (s, 3H) |
| රි | 8.34 (d, J = 5.1 Hz, 1H), 7.24 (br s, 4H), 6.96 (d, J = 5.1 Hz, 1H), |
| | 4.33 (q, J = 8.6 Hz, 2H), 4.13 (m, 1H), 3.45 (m, 1H), 3.30 (m, 1H), |
| | 2.98 (m, 2H), 2.82 (m, 1H), 2.46 and 2.49 (s, 3H), 2.41 (m, 1H), 2.24 |
| | and 2.27 (s, 3H), 2.10 (m, 2H), 1.96 (m, 1H), 1.15-1.90 (m, 7H), 0.92 |
| | (s, 3H) |
| മ | 8.92 (s, 1H), 7.23 (br s, 4H), 4.11 (m, 1H), 3.79 (s, 3H), 3.30-3.45 (m, |
| | 2H), 2.97 (m, 2H), 2.81 (m, 1H), 2.45 and 2.42 (s, 6H), 2.40 (m, 1H), |
| | 1.90-2.20 (m, 3H), 1.15-1.90 (m, 7H), 0.92 (s, 3H) |

ഗ Example 7

Alternate synthesis of the compounds of Example 6.

The product of Example 6, step B-2 (566 mg, 1.20 mmol) is treated with H₃CONH₂·HCI using conditions similar to those shown in Example 6,

step B-3. The resulting crude mixture of Z- and E-methoximes is separated on a preparative silica gel TLC plate (eluting with hexanes/ EtOAc, 80:20) to afford, in order of elution, first the E-methoxime (175 mg; 29%), then the Z-methoxime (175 mg; 29%), both products as oils.

2) The Z-methoxime (75 mg; 0.15 mmol) of step 1 is deprotected following conditions similar to those shown in Example 6, step B-5 and the resulting free amine (46 mg) is directly subjected to amidation with 2,4-dimethylnicotinic acid using conditions similar to those shown in Example 6, step B-6 to yield 50 mg (82%) of a colorless oil.

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The following compounds are prepared via similar procedures:

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wherein R4, R6 and R2 are as defined in the table:

| 7G F3C- CH3- CI | 7F F ₃ CO- CH ₃ - H ₃ C | 7E F ₃ C- CH ₃ - H ₃ C | 7D F3CO- CH3CH2- H3C | 7C F ₃ CO- CH ₃ - H ₃ C | 7B F ₃ CO· CH ₃ · C | 7A F3CO- CH3- H3C | Ex. R ⁶ R ⁴ A |
|--------------------|--|---|-----------------------------------|--|---|-------------------|-------------------------------------|
| CH ₃ - | CH ₃ - | CH ₃ - | CH ₃ CH ₂ - | 어. | CH ₃ - | Сн ₃ - | 72 |
| CI NH ₂ | н₃СДон | н₃с-Дсн₃ | н₃СДСН₃ | H ₃ C NH ₂ | CI NIH ₂ | н₃с∰сн₃ | A2 |
| 537.2234 | 534.2571 | 516.2833 | 546.2940 | 533.2730 | 553.2192 | 532.2795 | HRMS (MH+) found |

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PCT/US00/11633

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| 7W | 7V | 7U (E isomer) | 71 | 78 | 7R | 70 | 7P (E isomer) | .70 | 7N | 7M | 71 | 7K | 7,1 | 71 | 7H |
|-----------------------------------|--------------------|---------------------|-----------------------------------|---|---|--------------------|---------------------|-----------------------------------|--------------------|--|-----------------------------------|-------------------|-----------------------------------|-----------------------------------|--------------------|
| F ₃ CO- | F ₃ CO- | F ₃ CO- | F ₃ C- | F ₃ C- | F ₃ CO- | F ₃ CO- | F ₃ C- | F ₃ CO- | F ₃ CO- | F ₃ CO- | F ₃ C- | F ₃ C- | F ₃ CO- | F ₃ C- | F ₃ C- |
| CH ₃ CH ₂ - | H ₃ CO | CH ₃ - | CH ₃ CH ₂ - | CH ₃ CH ₂ - | CH ₃ CH ₂ - | CH ₃ - | СН ₃ - | CH ₃ CH ₂ - | | CH ₃ CH ₂ - | CH ₃ CH ₂ - | СН3- | CH ₃ CH ₂ - | CH ₃ - | CH ₃ - |
| the Action | н₃СССН₃ | н₃С√Ссн₃ | HC CH | H _C C L _N CH _B | H3C \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ | HSC CH Mayo | њс√‱сњ | H ₃ C ₂ OH | н₃СССН₃ | н _я с- _С сн _я | нзсфон | ᠊ᡰᢑᡄᠸᢅᡳᡋᠲ | OL NH2 | H ₃ C √NH ₂ | CI NH ₂ |
| 563.2855 | 576.3049 | 532.2784 | 547.1348 | 531.1002 | 590.2854 | 549.2686 | 517.2831 | 548.2732 | 572.3093 | 547.2888 | 532.2787 | 517.2812 | 567.2362 | 537.2234 | 537.2234 |

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| о 577.3019 нс сь 577.3019 рle 7: | F | Nam | 7HH F ₃ CO- CH ₃ (CH ₂) ₂ - H _C Additional data for compounds of Example 7: | F3CO- ional data for | 7HH Addition |
|--|----------|--|---|-------------------------|-----------------|
| 18 | 617.1918 | , CO | CH ₃ (CH ₂) ₂ - | F ₃ CO- | 766 |
| 30 | 562.3030 | H ₃ C N CH ₃ | CH ₃ (CH ₂) ₂ - | F ₃ CO- | 7FF |
| 48 | 603,1748 | 0+z | CH3CH2- | F ₃ CO- | 7EE |
| 6 | 589.1610 | 0+V C | CH ₃ - | F ₃ CO- | 700 |
| - 61 | 548.2861 | H ₃ C N N N N N N | CH ₃ CH ₂ - | F ₃ CO- | 700 |
| 21 | 587.1821 | CI N | CH ₃ CH ₂ - | F3CO- | 788 |
| .38 | 573.1638 | Z | сн _з - | F3CO- | 7AA |
| 37 | 641.1537 | × | CF3CH2- | F3CO- | 77 |
| 89 | 574.2889 | но∰он | | F ₃ CO- | 74 |
| 52 | 573.3052 | 45 A C+ | | F ₃ CO- | 7× |

| | | 000. | 5 - - | |
|---|----|----------|--------------------|----|
| | _[| 589 1610 | ¥. | ╝ |
| (d. 3H), 1.45-2.20 (m. 9H), 1.20 (t. 3H), 0.90 (s. 3H) | | | Z | |
| 4.05 (q, 2H), 3.20-3.50 (m, 2H), 2.80-3.15 (m, 3H), 2.45 (d, 3H), 2.25 | | | H3C4 CH3 | |
| 8.31 (d, 1H), 7.61 (d, 2H), 7.32 (d, 2H), 6.95 (d, 2H), 4.25 (m, 2H), | 78 | 548 2861 | -\$ 2 | |
| 7H), 1.28 (t, J = 7.1 Hz, 3H), 0.88 (br s, 3H) | | | <u>/</u> | |
| 2.39 (m, 1H), 2.15-2.30 (m, 6H), 1.85-2.15 (m, 3H), 1.10-1.85 (m, | | 587.1821 | <u>δ</u> | |
| J = 7.1 Hz, 2H), 3.15-3.50 (m, 2H), 2.86-3.10 (m, 2H), 2.80 (m, 1H), | | | Z | ┸ |
| 7.13-7.30 (m, 5H), 7.14 (m, 1H), 6.95 (m, 1H), 4.13 (m, 1H), 4.03 (q, | 7R | | | _ |
| (m, 3H), 1.15-1.85 (m, 7H), 0.88 (s, 3H) | | 573.1638 | -{ | |
| J = 11.6 Hz, 3H), 2.45 (m, 1H), 2.20 (d, J = 11.6 Hz, 3H), 1.85-2.20 | | | <u> </u> | |
| (d, | | | 5 2 2 | |
| 4.16 (m, 1H), 3.75 (s, 3H), 3.20-3.45 (m, 2H), 2.85-3.00 (m, 3H), 2.41 | | 641 1537 | 3 6 | 丄 |
| 7Q $8.11 (d, J = 6.8 Hz, 1H), 7.25 (br s, 4H), 6.94 (d, J = 6.8 Hz, 1H),$ | 70 | | _{ ((| |
| (d, 3H), 1.45-2.20 (m, 11H), 0.92 (s, 3H) | | 574.2889 | H.C. X | |
| 3.80 (3, 2H), 3.20-3.50 (m, 2H), 2.75-3.05 (m, 3H), 2.45 (d, 3H), 2.25 | | | Ç | L_ |
| 7K 8.31 (d, 1H), 7.61 (d, 2H), 7.31 (d, 2H), 6.95 (d, 2H), 4.30 (m, 2H), | | 5/3.3052 | 45 Act | |

Example 8

- თ and concentrated to yield 0.437 g (97%) of sulfide. 1.07 mmol) in DMF (25 ml) is added sodium methylmercaptide (0.113 g. mixture is then cooled to RT, diluted with Et₂O, washed with brine, dried 1.62 mmol) and the mixture is heated to 70° C for 12 h. The reaction To a stirred solution of the product of Example 5, step 8 (0.500 g,
- A solution of the product of step 1 (1.00 g; 2.31 mmol)
- ಠ H₃CONH₂·HCI (3.80 g, 46.2 mmol), and AcONa (3.79 g, 46.2 mmol) in subjected to flash chromatography over silica gel, to afford first the E-oxime CH₂Cl₂. The residue obtained after evaporation of the solvents is solvent, the residue is taken up in aqueous 0.1 N NaOH and extracted with EtOH (30 ml) is heated at reflux under N2 for 4 h. After evaporation of the
- ᆳ (eluting Et₂O/CH₂Ct₂, 1:4; 0.45 g; 24%), then the Z-oxime (0.25 g, 15%). the mixture is stirred at 0°C for 4 h. The reaction is then quenched with (5 ml) is at 0° C is added oxone (1.00 g, 1.627 mmol in 5 ml of CH₃OH) and To a solution of Z-oxime (0.250 g, 0.543 mmol) of step 2 in CH₃OH

7.55 (d, 2H), 7.30 (d, 2H), 7.15 (t, 1H), 6.75 (d, 1H), 6.60 (d, 1H),

4.25 (m, 2H), 3.80 (s, 3H), 3.40 (m, 2H), 2.80-3.20 (m, 3H), 2.40 (m,

1H), 1.40-2.20 (m, 13H), 0.90 (s, 3H)

1H-NMR (300 MHz 1H NMR (CDCI3))

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10% NaOH, concentrated, poured into water (10 ml) and extracted with CH₂Cl₂, dried and concentrated to yield 0.220 g (82%) of sulfone.

- 4) To a stirred solution of the product of step 3 (0.300 g, 0.608 mmol) in CH₂Cl₂ (5 ml) is added TFA(1 ml) and the mixture is stirred at RT for 2 h.
- 5 The reaction mixture is concentrated, poured into 10% NaOH and extracted with CH₂Cl₂. The combined extracts are dried and concentrated to afford 0.240 g (100%) of amine.
 5) To stirred solution of the product of step 4 (0.45 g, 0.114 mmol) in
- 5) To stirred solution of the product of step 4 (0.45 g, 0.114 mmol) in CH₂Cl₂ is added 2.6-dimethylnicotinic acid (0.26 g, 0.172 mmol), DEC
- 10 (0.33 g, 0.172 mmol), N,N,N-diisopropylethylamine (DIPEA) (0.2 ml) and HOBT (0.24g, 0.172 mmol) and the mixture is stirred for 12 h at RT The reaction mixture is quenched with NaHCO₃, extracted with CH₂Cl₂, dried, concentrated and purified by preparative chromatography (20% EtOH/EtOAc) to afford 0.046 g (76%) of Z-oxime amide.
- 15 300 MHz ¹H NMR (CDCl₃) δ 8.32 (d, 1H), 7.95 (d, 2H), 7.40 (d, 2H), 6.95 (d, 1H), 4.20 (m, 1H), 3.82 (s, 3H), 3.30-3.45 (m, 3H), 3.10 (s, 3H), 2.80-3.00 (m, 3H), 2.50 (d, 2H), 2.25 (d, 2H), 1.30-2.20 (m, 12H), 0.92 (s, 3H). The following compounds were prepared in a similar manner:

20 wherein X, R⁶ and R² are as defined in the table:

| - 1 | | | · | | | | ٦. |
|-----|--|-------------------|--------------------|--------------------|---------------------------------|---------------------|--|
| | 8 E | 86 | 80 | 88 | 8A (mixture E/Z) | Ř | Wildigit X |
| | 0, KH3 | 6. 5. 7. 5. 7. | Br | 0, CH3 | 0 % CH ₃ | R | יון מווט דר מ |
| | CH ₃ CH ₂ Q -C- | CH3CH2Q | - CH30 | | -C- -C- -C-H ₃ | × | Wilelen A, it and it ale as delilled in the table: |
| | H ₃ C N-Y OH ₃ | Н3С√СН3 | CI NH ₂ | CI NH ₂ | Н3С СН3 | P2 | acie: |
| | 557.2798 | 541.2849 | 549.2133 | 547.2135 | 526.2753 | HRMS (MH+) found | |

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WO 00/66559 PCT/US00/11633

- 56 -

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8F 05 CH3 CH30 H3C CH3 543.2641

8G 05 CH3 CH30 H3C CH3 527.2692

8H F3C- CH30H20 H3C CH3 532.2895

8I 05 CH3 CH30 H3C CH3 532.2895

8 5 5 S each of the following solvents: THF, CH₂Cl₂ and CH₃OH, then wash with cartridge. To this mixture add 0.12 ml of a 1M solution of 5-methyl-3-[2of a 0.5M solution in DMF) and DMAP (25 μ l of a 0.05M solution in DMF). chlorophenyl]isoxazole-4-carboxylic acid in DMF (0.12 mmol), HOBT (86 µl slurry of 0.25 g (~ 0.22 mmol) of resin bound cardodlimide (prepared by 572 (calculated MW 571); TLC R_f = 0.45 (CH₂Cl₂/CH₃OH/ NH₄OH reduced pressure to afford the title compound. LCMS found MH*= 570 Shake this mixture for 14 h, filter and add 0.3 g of Amberlyst-15 resin (~ 1.5 carbodiimide in DMF at 100 C) in DMF (2 ml) in a polyethylene SPE Stock solution A \sim 0.1M). Add 430 μ l of stock solution A (0.043 mmol) to a min, and 1 time for 5 min.). Combine and concentrate the filtrates under THF and CH₂Cl₂. Treat the resin with 2M NH₃ in CH₃OH (1 time for 30 mmol) to the filtrate. Shake for 1 to 2 h, filter and wash the resin twice with reacting Argopore-Cl resin with 1-(3-dimethylaminopropyl)3-ethyl Dissolve the starting amine (2.0 g, 5.7 mmol) in CHCl₃ (57 ml; =

wherein R2 is as defined in the table:

| - 1 | | | | | , | | |
|-----|---------------------------------|-------|--|-----------------------|--|---|------------------------------|
| | | 96 | 90 | 90 | 98 | 9A | , m |
| | Š | abla | Ğ | H ₃ C N | H ₃ C-\NH ₂ | H3C | R ² |
| | MH+ = 497.1 $R_t = 6.32 min$ | LCMS: | LCMS: MH+ = 507.1 $R_1 = 6.39 \text{ min}$ | LCMS: MH+ = 606 | MS m/e = 475.2, 477.2 (Electrospray) | LCMS: MH+ = 538.1 R _t = 6.27 min | · Data |
| | 0.48 | | 0.49 | 0.57 | | 0.58 | TLC R _f values |

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chromatography over silica gel (eluting CH₂Cl₂/CH₃OH 97:3 to 95:5) to into 5% aqueous NaHCO3, extracted with CH2Cl2, and dried over Na2SO4. After concentration of the solvents, the resulting oil was purified by flash (2 ml) at 0 °C was added diethylazodicarboxylate (160 ml; 1 mmol) and the pyridine (95.1 mg; 1 mmol) and PPh₃ (262 mg; 1 mmol) in anhydrous THF mixture was allowed to warm to RT overnight. The reaction was poured Step 1: To a solution of alcohol 39ab (406 mg; 0.87 mmol), 3-hydroxy-

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mg; 0.53 mmol) proceeded as in Example 2 to obtain the desired amine afford the desired compound (290 mg; 61%), as an oil. (210 mg; 89%), as a white foam. Step 2: Removal of the Boc-protecting group of the product of step 1 (290

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WO 00/66559 PCT/US00/11633

ഗ Hz, 1H), 7.43 (br d, J = 8.4 Hz, 2H), 7.14 (br d, J = 8.4 Hz, 2H), 6.95-7.10 (m, 2H), 4.75 (br d, J = 6.8 Hz, 1H), 4.15 (m, 1H), 3.44 (m, 1H), 3.33 (m, Example 2 to obtain the title compound (32 mg; 49%) as a colorless oil: dimethylpyrimidine-5-carboxylic acid following the conditions described in Step 3: The amine of step 2 (50 mg; 0.11 mmol) was coupled with 4,6-1.65-1.85 (m, 2H), 1.15-1.50 (m, 5H), 0.90 (s, 3H); HRMS (MH+) 1H), 2.95 (m, 2H), 2.79 (m, 1H), 2.42 and 2.44 (s, 3H), 1.85-2.15 (m, 3H), ¹H-NMR (300 MHz, CDCl₃) δ 8.91 (s, 1H), 8.20 (br s, 1H), 8.10 (d, J = 4.5

5 prepared Using similar procedures, compounds of the following structure were

wherein R3 R6 and R2 are as

| 567.2308 | H ₃ C N N N CH ₃ CH ₃ | , <u> </u> | Ω | 109 |
|---------------------|---|------------|---------|--------|
| 595.2072 | H3C√KCH3 N≪N | | Br | 10G |
| 532.2981 | 40 T CH | ţ.© | 711 | 10F |
| 516.3031 | Н₃С√СН₃ | Į. | 711 | 106 |
| 517.2992 | H₃C√~CH₃ N≪N | Į.© | П | 100 |
| 595.2078 | H ₃ C N N N N N | ئي الم | Br | 100 |
| 577.2166 | НэС√СН3 | (کی) | Br | 108 |
| 592.2848 | HC Toth | Ţ) | снзѕо2- | 10A |
| HRMS (MH+) found | R2 | Ą | 공 | , X |

| ហ | | | | | | | | • |
|---|--|--|---|---|---|--|--|---|
| o. | | 10L | | OR . | 10H | Į. | | Additi 10C |
| Example 11 CH ₃ H ₃ C N CH ₃ H ₃ C O CH ₃ PhB(OH) ₂ PhB(OH) ₃ PhB(OH) ₂ PhB(OH) ₂ PhB(OH) ₃ PhB(O | 3.30 (m, 1H), 3.02 (s, 3H), 2.99 (m, 2H), 2.79 (m, 1H), 2.47 and 2.48 (s, 3H), 2.45 (m, 1H), 2.25 and 2.26 (s, 3H), 1.65-2.15 (m, 5H), 1.15-1.55 (m, 5H), 0.90 (s, 3H) | 8.33 (d, J = 5.1 Hz, 1H), 7.99 (dd, J = 4.8 and 1.8 Hz, 1H), 7.86 (d, J = 8.4 Hz, 2H), 7.56 (d, J = 8.4 Hz, 2H), 7.53 (m, 1H), 6.96 (d, J = 8.4 Hz, 2H), 7.53 (m, 1H), 6.96 (d, J = 8.4 Hz, 2H), 7.53 (m, 1H), 6.96 (d, J = 8.4 Hz, 2H), 7.53 (m, 1H), 6.96 (d, J = 8.4 Hz, 2H), 7.53 (m, 1H), 6.96 (d, J = 8.4 Hz, 1H), 6.96 (d, J = 8.4 Hz, 2H), 7.53 (m, 1H), 7.53 (m, 1H | Hz, 2H), 6.90 (m, 1H), 6.74 (m, 1H), 6.59 (m, 1H), 4.83 (d, <i>J</i> = 6.8 Hz, 1H), 4.08 (m, 1H), 3.20-3.40 (m, 2H), 2.70-3.00 (m, 3H), 2.35 (br s, 3H), 1.65-2.15 (m, 5H), 1.15-1.50 (m, 5H), 0.87 (s, 3H) | (br d, $J = 6.8$ Hz, 1H), 4.19 (m, 1H), 3.46 (m, 1H), 3.37 (m, 1H), 3.00 (m, 2H), 2.81 (m, 1H), 2.47 and 2.49 (s, 3H), 1.90-2.15 (m, 3H), 1.65-1.90 (m, 2H), 1.20-1.50 (m, 5H), 0.93 (s, 3H) 8.81 (s. 1H), 7.78 (d. $J = 8.4$ Hz, 2H), 7.53 (m. 1H), 7.47 (d. $J = 8.4$ | 8.95 (s, 1H), 7.32 (br d, J = 8.4 Hz, 2H), 7.23 (br d, J = 8.4 Hz, 2H), 7.08 (t, J = 8.1 Hz, 1H), 6.80-6.90 (m, 2H), 6.68 (m, 1H), 4.77 | 8.17 (a, $J = 6.8$ Hz, 1H), 7.28 (m, 2H), 7.18 (t, $J = 7.5$ Hz, 1H), 6.95-7.10 (m, 3H), 6.87 (t, $J = 7.5$ Hz, 1H), 6.80 (d, $J = 7.5$ Hz, 2H), 4.80 (d, $J = 6.8$ Hz, 1H), 4.17 (m, 1H), 3.25-3.50 (m, 2H), 2.99 (m, 2H), 2.80 (m, 1H), 2.43 (br s, 3H), 2.24 (br s, 3H), 1.65-2.20 (m, 5H), 1.15-1.50 (m, 5H), 0.90 (s, 3H) | (m, 2H), 1.20-1.50 (m, 5H), 0.93 (s, 3H) | Additional data for compounds of Example 10: Ex. 1H-NMR (300 MHz ¹ H NMR (CDCl ₃)) 10C 8.95 (s, 1H), 7.46 (br d, J = 8.4 Hz, 2H), 7.17 (br d, J = 8.4 Hz, 2H) |

- 59 -

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100

F₃CO-

584.2860

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F₃C-

585.2559

10S

F₃C

H3C/ CH3

601.2556

10R

F₃CO-

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598.2903

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F₃CO-

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583.2905

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H₃C N CH₃

597.2951

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CH3SO2-

H₃C N CH₃

595.2749

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CH3SO2-

H₃C N CH₃

611.2460

H₃C N N N N N

617.2514

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F₃CO-

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F₃CO-

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601.2817

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СH₃SO₂-

H₃C N CH₃

595.2764

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CH3SO2-

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577.2853

582.2955

<u>5</u>

F₃C-

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N-Boc-4-piperidone (10 g, 50 mmol) and PPh₃ (53 g, 200 mmol) were taken up in CH₃CN (100 ml). The solution was cooled to 0 °C and CBr₄ (33 g, 100 mmol) was added to the solution at 0 °C. The solution was stirred at 0 °C for 15 min. and at 25 °C for 2 h. Et₂O (200 ml) was added,

and the resulting mixture was filtered through a plug of SiO₂.

Concentration gave a yellow solid. Purification via flash chromatography (9/1 hexanes/Et₂O, SiO₂) gave 10 g (56 %) of the di-bromo product as a

10 2) A solution of the product of step 1 (1 g, 2.8 mmol), PhB(OH)₂ (1.2 g, 9.9 mmol), PdCl₂(PPh₃)₂ (197 mg, 0.28 mmol), and Na₂CO₃ (897 mg, 8.5 mmol) were taken up in THF/H₂O (4/1, 20 ml) and stirred at 65 °C under N₂ for 24 h. The solution was partitioned between EtOAc and H₂O, the aqueous layer was extracted with EtOAc and the combined organic layers

were washed with brine and dried over Na₂SO₄. Filtration and concentration gave a dark brown oil. Purification via flash chromatography (9/1 hexanes/El₂O, SiO₂) gave 941 mg (96 %) of the desired product as a white solid, m.p. = 152-153 °C.

3) A solution of the product of step 2 (500 mg, 1.4 mmot) and Pd(OH)₂ on carbon (100 mg, 20 wt % Pd (dry basis), 50 wt % H₂O) were taken up in CH₃OH (20 ml) and shaken in a Parr apparatus under H₂ (50 psi) for 15 h. The mixture was filtered and concentrated to give 501 mg (99 %) of the diphenylmethyl piperidine as a colorless oil.

4) TFA (1.4 ml) was added to a solution of the product of step 3 (500 mg, 1.4 mmol) in CH₂Cl₂ (15 ml). The solution was stirred at 25 °C for 23 h. The solution was concentrated and the residue partitioned between CH₂Cl₂ and 1 N NaOH. The aqueous layer was extracted with CH₂Cl₂, the combined organic layers were dried over Na₂SO₄, filtered and concentrated to obtain 349 mg (99 %) of the free amine as a yellow oil,

WO 00/66559 PCT/US00/11633

- 62 -

m.p. (HCl) = decomp. above 220-230 °C. HRMS calc'd for C₁₈H₂₂N (MH³): 252.1752, Found: 252.1751.

5) A solution of the product of step 4 (349 mg, 1.4 mmol), N-Boc-4-piperidone 280 mg, 1.4 mmol), and Ti(OiPr)₄ (0.42 ml, 1.4 mmol) were
 5 taken up CH₂Cl₂ (15 ml) under N₂. After stirring at 25 °C for 17 h, Et₂AlCN (2.8 mmol, 2.8 ml of 1.0 M in toluene) was added and the solution was

stirred an additional 18 h at 25 °C. The solution was quenched with sat

- NaHCO₃, diluted with EtOAc and filtered through Celite. The aqueous layer was extracted with EtOAc and the combined EtOAc layers were dried over Na₂SO₄. Filtration and concentration gave a yellow oil. Purification via preparative layer chromatography (3/1 hexanes/EtOAc, SiO₂) gave 430 mg (67 %) of the desired product as an oil.
- 6) A solution of the product of step 5 (430 mg, 0.94 mmol) in THF (20 ml) was cooled to 0 °C under N₂. CH₃MgBr (1.6 ml of 3.0 M in Et₂O, 4.7
- 15 mmol) was added at 0 °C and the solution stirred at 25 °C for 19 h. The reaction mixture was quenched with sat. NH₄Cl, diluted with CH₂Cl₂ and 1 N NaOH (check aqueous layer with pH paper, pH = 8-10). The layers were separated and the aqueous layer extracted with CH₂Cl₂. The combined organic layers were dried over Na₂SO₄, filtered and concentrated to obtain a yellow oil. Purification via flash chromatography (3/1 hexanes/EtOAc,
- SiO₂) gave 275 mg (65 %) of the product as a yellow oil.

 7) TFA (0.60 ml) was added to a solution of the product of step 6 (275 mg, 0.61 mmol) in CH₂Cl₂ (15 ml) and the solution was stirred at 25 °C for
- 18 h. The solution was concentrated and the residue was partitioned between CH₂Cl₂ and 1 N NaOH. The aqueous layer was extracted with CH₂Cl₂, the combined organic layers were dried over Na₂SO₄, filtered and concentrated to obtain 209 mg (99 %) of thje amine as a yellow oil. HRMS calc'd for C₂₄H₃₃N₂ (MH*): 349.2644, Found: 349.2638.
- 8) A solution of the product of step 7 (50 mg, 0.14 mmol), 2.6-dimethyl-30 benzoic acid (63 mg, 0.42 mmol), EDCI (54 mg, 0.28 mmol), HOBT (38 mg, 0.28 mmol), and iPr₂NEt (0.10 ml) were taken up in CH₂Cl₂ (3 ml). The solution was stirred at 25 °C for 18 h, then diluted with CH₂Cl₂ and washed with 1 N NaOH. The aqueous layer was extracted with CH₂Cl₂, the combined organic layers were dried over Na₂SO₄, and filtered and
- 35 concentrated to give a yellow oil. Purification via preparative thin-layer chromatography (3/1 hexanes/EtOAc SiO₂) gave 47 mg (70 %) of the title

for C₃₃H₄₁N₂O (MH⁺): 481.3219, Found: 481.3225 compound as a colorless oil, m.p. (HCl salt) = 195-201 °C. HRMS calc'd

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prepared Using similar procedures, compounds of the following structure were

wherein R6 and R2 are as defined in the table

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| 1F | Ħ | 10 | 10 | 18 | 1A | Ϋ́ |
|----------|----------------------------------|--------------------------------------|----------------------------------|-----------------------|----------------|------------------------|
| F,CO- | F,CO. | F,CO- | I | F,CO- | I | ЯG |
| н₃с√‱сн₃ | H ₃ C CH ₃ | H ₃ C N N N N | H ₃ C NH ₂ | н _э с-Донз | H₃C√ÃγCH₃ N | R² |
| 566.3020 | 582.2966 | 567.2957 | 482.3168 | 565.3069 | 482.3156 | HRMS (MH+) found |
| 175-181 | 92-98 | 175-181 | 187-192 | 204-209 | 201-207 | M.p., °C (HCi salt) |

Example 12

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WO 00/66559 PCT/US00/11633

- 64 -

- 60 mmol, 60 wt % in oil dispersion) was added to the solution at 25 °C. (12.6g, 55 mmol) were taken up in dry THF (50 ml) under N_2 . NaH (2.4 g. N-Boc-4-piperidone (10 g, 50 mmol) and diethyl benzylphosponate
- oil. Purification via flash chromatography (10/1 hexanes/Et₂O, SiO₂) gave brine and dried over MgSO₄. Filtration and concentration afforded a yellow extracted with EtOAc and the combined EtOAc layers were washed with partitioned between EtOAc and saturated NH₄Cl, the aqueous layer was The resulting mixture was heated at reflux for 3.5 h. The solution was
- ಠ 9.85 g (72 %) of the desired compound as a solid, m.p. = 63-65 °C. concentrated under reduced pressure. The crude product was taken up in mmol) at 0 °C. The solution was stirred at 0 °C for 15 min, then dropwise to a CH₂Cl₂ (100 ml) solution of the product of step 1 (5.0 g, 18 Bromine (1 ml, 20 mmol; dissolved in 10 ml CH₂Cl₂) was added
- 8 5 Purification via flash chromatography (7/1 hexanes/Et₂O, SiO₂) gave 5.2 g dried over MgSO₄. Filtration and concentration gave a yellow solid. then concentrated under reduced pressure. The residue was partitioned the solution in portions. The yellow mixture was stirred at 25 °C for 5h, tert-butanoi/THF (4/1, 100 ml), and KOtBu (4.1 g, 36 mmol) was added to with EtOAc, and the combined EtOAc layers were washed with brine and between EtOAc and saturated NH₄CI, the aqueous layer was extracted

(81 %) of the desired product as a yellow solid. m.p. = 80-83 °C

- organic layers were dried over Na₂SO₄, filtered and concentrated to obtain above 185-195 °C. HRMS calc'd for C12H15BrN (MH*): 254.0367, Found: 1.46 g (98 %) of the amine as an orange oil, m.p. (HCl salt) = decomp. concentrated and the residue was partitioned between CH₂Cl₂ and 1 N 5.9 mmol) in CH₂Cl₂ (25 ml). The solution was stirred at 25 °C for 5 h, NaOH. The aqueous layer was extracted with CH2Cl2 and the combined TFA (5.9 ml) was added to a solution of the product of step 2 (2.1 g
- ಠ up in CH2Cl2 (30 ml) under N2. After stirring at 25 °C for 18 h, Et2ALCN piperidone (1.1 g, 5.6 mmol), and Ti(OiPr)₄ (1.7 ml, 5.6 mmol) were taken quenched with sat. NaHCO3, diluted with EtOAc and filtered through Celite solution was stirred an additional 18 h at 25 °C. The solution was (6.7 mmol, 6.7 ml, 1.0 M in toluene) was added to the solution and the A solution of the product of step 3 (1.4 g, 5.6 mmol), N-Boc-4-
- ឆ 2.0 g (78 %) of the desired product as an off-white solid. oil. Purification via flash chromatography (3/1 hexanes/EtOAc, SiO₂) gave layers were dried over Na₂SO₄. Filtration and concentration gave a yellow The aqueous layer was extracted with EtOAc and the combined EtOAc
- 8 stirred at that temperature for 16 h. The reaction mixture was quenched was cooled to 0 °C under N2. CH3MgBr (7.2 ml of 3.0 M in Et2O, 21 mmol with sat. NH₄Cl and diluted with CH₂Cl₂ and 1 N NaOH (check aqueous was added to the solution at 0 °C. The solution was warmed to 25 °C and A solution of the product of step 4 (2.0 g, 4.3 mmol) in THF (30 ml)
- 23 g (82 %) of the desired product as a yellow oil. dried over Na₂SO₄. Filtration and concentration gave a yellow oil. layer was extracted with CH₂Cl₂ and the combined organic layers were Purification via flash chromatography (3/1 hexanes/EtOAc, SiO₂) gave 1.56 layer with pH paper, pH = 8-10). The layers were separated, the aqueous
- မွ stirred at 65 °C under N₂ for 18 h. The solution was partitioned between 4-CF₃C₆H₄B(OH)₂ (380 mg, 2 mmol), PdCl₂(PPh₃)₂ (50 mg, 0.067 mmol) combined organic layers were washed with brine and dried over Na₂SO₄ EtOAc and H₂O and the aqueous layer was extracted with EtOAc. The and Na₂CO₃ (210 mg, 2 mmol) were taken up THF/H₂O (4/1, 15 ml) and A solution of the product of step 5 (300 mg, 0.67 mmol),
- 35 chromatography (4/1 hexanes/EtOAc, SiO2) gave 229 mg (67 %) of the Filtration and concentration gave a dark brown oil. Purification via flash desired product as a colorless oil.

WO 00/66559

PCT/US00/11633

- G (100 %) of the (±)-product as a colorless foam. HRMS calc'd for taken up in CH₃OH (35 ml) and shaken in a Parr apparatus under H₂ (50 $Pd(OH)_2$ on carbon (200 mg, 20 wt % Pd (dry basis), 50 wt % H_2O) were C₃₀H₄₀O₂N₃ (MH*): 517.3042, Found: 517.3050. psi) for 20 h. The mixture was filtered and concentrated to obtain 232 mg A solution of the product of step 6 (229 mg, 0.45 mmol) and
- ಕ organic layers were dried over Na₂SO₄, filtered and concentrated to obtain 1 N NaOH. The aqueous layer was extracted with CH2Cl2, the combined mg, 0.45 mmol) in CH₂Cl₂ (15 ml). The solution was stirred at 25 °C for 24 h, then concentrated and the residue was partitioned between CH₂Cl₂ and 146 mg (78 %) of the (±)-amine as a yellow oil. TFA (0.45 ml) was added to a solution of the product of step 7 (235
- 8 5 preparative thin-layer chromatography (1/1 acetone/hexanes SiO₂) gave 0.50 mmol), HOBT (70 mg, 0.50 mmol), and iPr₂NEt (0.17 ml) was taken with CH₂Cl₂ and washed with 1 N NaOH. The aqueous layer was up in CH₂Cl₂ (3 ml). The solution was stirred at 25 °C for 18 h, then diluted dimethylpyrimidine-5-carboxylic acid (110 mg, 0.75 mmol), EDCI (96 mg 186-191 °C. HRMS calc'd for C₃₂H₃₈N₄OF₃ (MH⁺): 551.2998, Found: Na₂SO₄, filtered and concentrated to obtain a yellow oil. Purification via extracted with CH₂Cl₂, the combined organic layers were dried over 121 mg (88 %) of the title compound as a colorless oil, m.p. (HCl salt) = A solution of the product of step 8 (102 mg, 0.25 mmol), 4,6-

made by the following process: The 4,6-dimethylpyrimidine-5-carboxylic acid used in step 9 was

trifluoromethane sulfonate (88.6 g) was added dropwise and after addition ml) were mixed together, using an overhead mechanical stirrer. CH₃CN Step 1: Ethyl diacetoacetate (93.4 g), Cs₂CO₃ (185 g) and CH₃CN (550 (50 ml) was added and the resulting mixture was cooled to 0°C. Methyl

- ႘ၟ မ Et₂O extracts were combined and evaporated to half volume. The solution extracts were combined and Et₂O (300 ml) was added. The resulting filtered, and the salts were washed with Et $_2$ O (2 X 50 ml). The organic mixture was filtered, the filter cake was washed with El₂O (2 X 100 ml), the the cooling bath was removed. The mixture was stirred for 1 h at RT,
- was cooled in an ice bath and washed once with cooled (0°C) 2 N NaOH

to give the desired product as a yellow liquid (64.7 g) in 65% yield, which was used directly in the next step. (pH = 11). The Et₂O layer was dried over MgSO₄, filtered and evaporated

G removed under vacuum. The resulting liquid was partitioned between to RT, the resulting precipitate was filtered off and the ethanol was were mixed together at RT. After refluxing for 4 h, the mixture was cooled (commercial solution; 21 wt%; 113 g) and formamidine acetate (36.2 g) Step 2: The product of step 1 (64.2 g), sodium ethoxide in ethanol water and CH₂Cl₂ and the aqueous layer was extracted with CH₂Cl₂ (3 \times

- **5** evaporation of the appropriate fractions, the desired product (28.5 g) was isotated in 46% yield and used directly in the next step. gel chromatography (980 g; 4:1 hexanes:EtOAc as eluant). After evaporated to give a dark crude liquid (50.7 g) which was purified by silica 150 ml). The CH₂Cl₂ extracts were dried over MgSO₄, filtered and
- ᇙ in vacuo until a thick paste resulted. Water (20 ml) was added, the mixture EtOH (130 ml) were mixed together at RT and heated at reflux for 1h. The was cooled to 0°C and conc. HCl (14.3 ml) was added dropwise with resulting solution was cooled to RT and the volatile materials were removed Step 3: The product of step 2 (28.1 g), NaOH (6.72 g), water (65 ml) and
- 8 stirring. The resulting white precipitate was collected by filtration, washed removed in vacuo at 50°C and then dried under vacuum (1 mm Hg) for 18 resulting white solid was treated with toluene $(2 \times 20 \text{ m})$, the solvent was with ice water (2 X 10 ml) and air dried with suction for 30 min. The The desired product (14.9 g) was isolated as a white solid in 63% yield
- 25 5.30%, N 18.41%; found: C 55.13%, H 5.44%, N 18.18% mp: 176-178°C. Elemental analysis of C₇H₈N₂O₂: calc'd C 55.26%, H

product (4.68 g) as a cream colored solid to give a combined yield of 83% washed with ice water (2 X 5 ml) and dried as described above to give the precipitate formed was collected by filtration. The resulting solid was resulting mixture was stirred at RT for 5 min, cooled in an ice bath and the aqueous filtrate (from above) to dryness and addition of water (20 ml). The A second crop of product was isolated by evaporation of the

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WO 00/66559 PCT/US00/11633

ಕ G saturated NaHCO₃ then brine. The residue obtained after concentration of the organic layer is purified by flash chromatography over silica gel (eluting to 0 °C, stirred 30 min at this temperature, and a solution of the product of 4.80 mmol) in anhydrous THF (15 ml) at -40 °C is added n-BuLi 2.5 N in allowed to warm to RT overnight, poured into CH₂Cl₂, and washed with hexanes (2.12 ml; 5.3 mmol) via syringe. The reaction is allowed to warm Step1: To a suspension of methyltriphenylphosphonium bromide (1.89 g: Example 6, step B-2 (2.24 g; 4.8 mmol) is added. The solution is then

5 of this solution (1.5 ml; 0.59 mmol of theoretical intermediate) is added to a 0.5 N in THF (3 ml; 1.5 mmol) is refluxed 2 h under inert atmosphere. Part Step 2: A solution of the product of step 1 (0.56 g; 1.2 mmol) and 9-BBN

with CH₂Cl₂/EtOAc, 9:1) to afford 0.56 g (25%) of an oil

and water (80 µl). The reaction is stirred 2 h at 60 °C and overnight at RT, poured into 5% aqueous NaHCO3, and extracted with CH2Cl2. Combined (19.8 mg), triphenylarsine (24.1 mg) and Cs₂CO₃ (250 mg) in DMF (0.40 ml mixture of 1-chloro-3-iodobenzene (88 μl; 0.71 mmol), PdCl₂dppf.CH₂Cl₂

8 chromatography over silica gel (eluting with EtOAc/hexanes, 8:2) to provide organic layers are dried over Na₂SO₄, concentrated, and purified by 100 mg (29%) of an oil.

25 86%). This amine (45 mg; 0.09 mmol) was coupled with 4,6-dimethylmmol) was removed as in Example 2 to obtain the desired amine (70 mg; Step 3: The Boc-protecting group of the product of step 2 (100 mg; 0.17

pyrimidine-5-carboxylic acid following the conditions described in Example

2 to obtain the title compound as a colorless oil (32 mg). 1H-NMR (300 MHz, CDCl₃) δ 8.93 (d, J = 3.8 Hz, 1H), 6.90-7.10 (m, 5H), 6.88 (br s, 1H), 6.71 (d, J = 7 Hz, 1H), 4.20 (m, 1H), 3.25-3.55 (m, 2H), 3.19 (m, 2H), 2.50 3.10 (m, 5H), 2.47 and 2.48 (s, 3H), 2.42 and 2.43 (s, 3H), 1.70-2.20 (m, 5H), 1.20-1.65 (m, 5H), 0.92 (s, 3H); HRMS (MH+) 615.2722.

Using a similar procedure, the following compound was also prepared: ທ

Example 14

Enantiomers I and II

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To prepare a compound wherein R2 is 2,6-dimethylphenyl:

A solution of the product of step 5 in example 12 (300 mg, 0.67 mmol), 4-CF₃OC₈H₄B(OH)₂ (410 mg, 2 mmol), PdCl₂(PPh₃)₂ (50 mg, 0.067

WO 00/66559 PCT/US00/11633

mmol), and Na,CO $_3$ (210 mg, 2 mmol) were taken up in THF/H $_2$ O (4/1, 15' ml) and stirred at 65 $^{\circ}$ C under N $_2$ for 19 h. The solution was partitioned

between EtOAc and H₂O, and the aqueous layer was extracted with EtOAc

- The combined organic layers were washed with brine and dried over 5 Na,SO₄. Filtration and concentration gave a dark brown oil. Purification via flash chromatography (4/1 hexanes/Et,O, SiO₅) gave 356 mg (100 %) of the desired product as a yellow oil.
- A solution of the product in step 1 (340 mg, 0.64 mmol) and Pd(OH), on carbon (300 mg, 20 wt % Pd (dry basis), 50 wt % H₂O) were taken up in
- 10 CH₃OH (35 ml) and shaken in a Parr apparatus under H₂ (50 psi) for 18 h. The mixture was filtered and concentrated to obtain 341 mg (100 %) of the product, (±)-1, as a colorless foam.
- The amine (±)-1 was resolved via chiral HPLC separation. The conditions are as follows: CHIRALCEL® OD" (5 cm x 30 cm); Hexane/ isopropyl alcohol/diethylamine 75/25/0.05) at 25° C; 254 nm detection. The retention times for peak 1, (+)-enantiomer, and peak 2, (-)-enantiomer were 3.8 and 4.9 minutes, respectively [CHIRALCEL® OD" (hexane/ethanol/diethylamine 90/10/0.1) 25° C at 254 nm]. Peak 1 and peak 2 are the first
- retention times for peak 1, (+)-enantiomer, and peak 2, (-)-enantiomer were 3.8 and 4.9 minutes, respectively [CHIRALCEL® OD" (hexane/ethanol/diethylamine 90/10/0.1) 25° C at 254 nm]. Peak 1 and peak 2 are the first and second eluting peaks from the column, respectively. The enantiomers (I and II) were deprotected (CH₂CI₂TFA), and the free amine was coupled to the 2,6-dimethylbenzoic acid using the conditions described in example 11, steps 7 and 8. The hydrochloride salts were obtained by taking the free base up in EtOAc and triturating with 1 M HCI in EtO.

Data for the above compounds, 14A and 14B, and for additional compounds made in a similar manner, are given in the following table. In each case, the enantiomer designator I is derived from (+)-1 and the enantiomer designated II is derived from (-)-1.

| | Ar | | |
|----------|------------|-----|------|
| - | Enantiomer | | F300 |
| 185-190 | m.p. (HCI) | | |
| 565.3042 | calc | НВМ | · |
| (J) | | S | |

14A

found 65.3050

| | | | | | _ | | | |
|----------|----------|------------------|------------|----------|----------|----------|----------|--------------|
| 141 | 141 | 14H | 14G | 14F | 14E | 14D | 140 | 14B |
| BHN C | | -\ -\ | ₽ - | | | N N | Nen | \(\) |
| = | = | = | . # | = | _ | II | - | II |
| 195-200 | 193-198 | 145-151 | 214-218 | 180-185 | 195-201 | 170-175 | 168-174 | 175-180 |
| 651.3522 | 615.3010 | 658.3257 | 581.2991 | 582.2944 | 582.2944 | 567.2947 | 567.2947 | 565.3042 |
| 651.3526 | 615.3016 | 658.3251 | 581.2984 | 582.2958 | 582.2944 | 567.2957 | 567.2951 | 565.3050 |
| | | | | | | _ | | |

WO 00/66559 PCT/US00/11633

-72-

1) TFA

2) EDC,HOBT, 2,6-dimethylbenzoic Ex. 15

The dibromo-olefin (3.55 g, 10 mmol) and TFA (10 ml) were taken

layer was extracted with CH2Cl2. The combined organic layers were dried to give the cyano-amine as described in Step 5 of Example 11. up in CH2Cl2 and stirred at 25 °C for 20 h. The solution was concentrated. piperdine as a colorless oil. The free piperdine (2.41 g, 9.45 mmol) was (Na₂SO₄). Filtration and concentration gave the 2.4 g (94 %) of the free The residue was partitioned between CH,Cl, and 1 N NaOH. The aqueous treated sequentially with (a) N-Boc-4-piperidone/Ti(OiPr),, and (b) Et,AICN

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- 5 5 gave a yellow oil. Purification via flash chromatography (6/1 hexanes/ C₁₆H₂₂O₂N₂Br , 387.1647; Found, 387.1638. bromide as a solid. m.p. (free base) 85-90 °C. HRMS (MH') calcd. for EtOAc, SiO,) gave 2.54 g (69 % from the free piperidine) of the vinyl were washed with brine and dried (Na₂SO₄). Filtration and concentration quenched with 1 N NaOH and EtOAc. The mixture was filtered (Celite) taken up in THF (30 ml) and stirred at 25 °C for 19 h. The solution was The aqueous layer was extracted with EtOAc, the combined EtOAc layers The product of Step 1 and MeMgBr (16 ml, 3.0 M in Et₂O) were
- 20 25 gave a yellow oil. Purification via flash chromatography (3/1 to 1/1 as an oil. HRMS (MH') calcd. for C₂₅H₃₀O₂N₂F₃, 453.2729; Found bath) for 21 hours. The solution was partitioned between EtOAc and H₂O. 1.56 mmol) were taken up in THF/H2O (4/1, 10 ml) and heated at 75 °C (oil mg, 1.8 mmol), PdCl₂(PPh₃)₂ (36 mg, 0.052 mmol), and Na₂CO₃ (165 mg, were washed with brine and dried (Na,SO,). Filtration and concentration hexanes/EtOAc, SiO₂) gave 210 mg (89 %) of the phenyl substitued olefin The aqueous layer was extracted with EtOAc, the combined EtOAc layers The product of Step 2 (200 mg, 0.52 mmol), 4-CF₃C₆H₄B(OH)₂ (344
- ၶ HRMS (MH') calcd. for C2,H2,ON2F3, 487.2936; Found, 487.2928 title compound as a yellow oil (37 mg, 55%). m.p. (HCl salt) 130-140 °C. Example 11. The reduced product was deprotected and coupled to 2,6dimethyl benzoic acid as described in Example 11, steps 7-8 to give the The product of Step 3 was hydrogenated as described in Step 3 of

Using a similar procedure, the following compound was prepared:

PCT/US00/11633

- 73 -

m.p. (HCl salt) 135-145°C. HRMS (MH') calcd. for C₂₉H₃Q₂N₂F₃ , 503.2885; Found, 503.2896.

5 The following assays can be used to determine the CCR5 inhibitory and antagonistic activity of the compounds of the invention. <u>CCR5 Membrane Binding Assay:</u>

A high throughput screen utilizing a CCR5 membrane binding assay identifies inhibitors of RANTES binding. This assay utilizes membranes

O prepared from NIH 3T3 cells expressing the human CCR5 champking recentor.

10 prepared from NiH 3T3 cells expressing the human CCR5 chemokine receptor which have the ability to bind to RANTES, a natural ligand for the receptor. Using a 96-well plate format, membrane preparations are incubated with 125I-RANTES in the presence or absence of compound for one hour. Compounds are serially diluted over a wide range of 0.001ug/ml to 1 ug/ml and tested in 15 triplicates. Reaction cocktails are harvested through glass fiber filters, and

triplicates. Reaction cocktails are harvested through glass fiber filters, and washed thoroughly. Total counts for replicates are averaged and data reported as the concentration required to inhibit 50 percent of total 125I-RANTES binding. Compounds with potent activity in the membrane binding assay are further characterized in seconday cell-based HIV-1 entry and replication assays.

HIV-1 Entry Assay:

Replication defective HIV-1 reporter virions are generated by cotransfection of a plasmid encoding the NL4-3 strain of HIV-1 (which has been modified by mutation of the envelope gene and introduction of a luciferase reporter plasmid) along with a plasmid encoding one of several HIV-1 envelope genes as described by Connor et al. Virology, 206 (1995), p. 935-944. Following transfection of the two plasmids by calcium phosphate precipitation, the viral supernatants are harvested on day 3 and a functional viral titer determined. These stocks are then used to infect U87 cells stably

23

30 expressing CD4 and the chemokine receptor CCR5 which have been preincubated with or without test compound. Infections are carried out for 2 hours at 37 °C, the cells washed and media replaced with fresh media containing compound. The cells are incubated for 3 days, lysed and luciferase

WO 00/66559 PCT/US00/11633

activity determined. Results are reported as the concentration of compound required to inhibit 50% of the luciferase activity in the control cultures.

HIV-1 Replication Assay:

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This assay uses primary peripheral blood mononuclear cells or the stable U87-CCR5 cell line to determine the effect of anti-CCR5 compounds to block infection of primary HIV-1 strains. The primary lymphocytes are purified from normal healthy donors and stimulated *in vitro* with PHA and IL-2 three days prior to infection. Using a 96-well plate format, cells are pretreated with drug for 1 hour at 37 °C and subsequently infected with an M-tropic HIV-1 isolates. Ecllowing infection the cells are washed to compose desidual incenture.

10 isolates. Following infection, the cells are washed to remove residual inoculum and cultured in the presence of compound for 4 days. Culture supernatants are harvested and viral replication measured by determination of viral p24 antigen concentration.

Calcium Flux Assay

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Cells expressing the HIV coreceptor CCR5 are loaded with calcium sensitive dyes prior to addition of compound or the natural CCR5 ligand. Compounds with agonist properties will induce a calcium flux signal in the cell, while CCR5 antagonists are identified as compounds which do not induce signaling by themselves but are capable of blocking signaling by the natural ligand RANTES.

GTPyS Binding Assay (secondary membrane binding assay):

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A GTP_YS binding assay measures receptor activation by CCR5 ligands. This assay measures the binding of ³⁵S labeled-GTP to receptor coupled G-proteins that occurs as a result of receptor activation by an appropriate ligand. In this assay, the CCR5 ligand, RANTES, is incubated with membranes from

In this assay, the CCR5 ligand, RANTES, is incubated with membranes from CCR5 expressing cells and binding to the receptor activation (or binding) is determined by assaying for bound ³⁵S label. The assay quantitatively determines if compounds exhibit agonist characteristics by inducing activation of the receptor or alternatively antagonist properties by measuring inhibition of RANTES binding in a competitive or non-competitive fashion.

Chemotaxis Assay:

The chemotaxis assay is a functional assay which characterizes the agonist vs. antagonist properties of the test compounds. The assay measures the ability of a non-adherent murine cell line expressing human CCR5 (BaF-550) to migrate across a membrane in response to either test compounds or natural ligands (i.e., RANTES, MIP-18). Cells migrate across the permeable membrane towards compounds with agonist activity.

Compounds that are antagonists not only fail to induce chemotaxis, but are

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44

- 76 -

PCT/US00/11633

WO 00/66559

also capable of inhibiting cell migration in response to known CCR5

inflammatory conditions has been reported in such publications as The role of CC chemokine receptors such as CCR-5 receptors in

- ഗ (psoriasis); Journal of Allergy & Clinical Immunology, 100 (6, Pt 2) (1997), Bheumatology, 1Z (4) (1999), p. 419-425 (rheumatoid arthritis); Clinical & Immunology Letters, 5Z, (1997), 117-120 (arthritis); Clinical & Experimental nternational Journal of Immunopharmacology, 20 (11) (1998), p. 661-7 Experimental Immunology, 117 (2) (1999), p.237-243 (atopic dermatitis).
- ಠ p. \$52-5 (asthma); and <u>Journal of Immunology</u>, <u>159</u> (6) (1997), p. 2962-72

0.1 to 500 nM, and most preferably 0.1 to 100 nM. The results for of the invention range in activity from a Ki of 0.1 to 2000 nM, with preferred compounds having a range of activity from 0.1 to 1000 nM, more preferably In the assay to determine inhibition of RANTES binding, compounds

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table, "Ex. No." stands for "Example Number" and "nM" stands for determine inhibition of RANTES binding are given in the table below. In the preferred and representative compounds of formulas I and II in the test to

| | | _ | | | | | | | _ | | | | | | | |
|-----|------|-----|-----|-----|-----|-----|-----|-----|-----|------|------|-----|-----|-----|----|--------------------------------------|
| 9D | 7 | 6V | 5AB | 52 | 50 | 5N | 5L | 4C | 48 | 211 | 28 | 2G | 2 | 1.J | 18 | Ex. No. |
| 588 | 62.5 | 0.8 | 0.1 | 0.3 | 0.4 | 1.7 | 7.9 | 0.5 | 0.5 | 0.58 | 17.9 | 1.8 | 9.6 | 1 | 14 | Ki (nM) Inhibition of RANTES binding |

ಕ various compositions may be found in A. Gennaro (ed.), Remington's dosage forms suitable for oral administration. Examples of the art, e.g. magnesium carbonate, magnesium stearate, talc, sugar or to about 95 percent active ingredient. Suitable solid carriers are known in suppositories. The powders and tablets may be comprised of from about 5 include powders, tablets, dispersible granules, capsules, cachets and acceptable carriers can be either solid or liquid. Solid form preparations antagonist compounds described by this invention, inert, pharmaceutically Easton, Pennsylvania. Pharmaceutical Sciences, 18th Edition, (1990), Mack Publishing Co. pharmaceutically acceptable carriers and methods of manufacture for lactose. Tablets, powders, cachets and capsules can be used as solid For preparing pharmaceutical compositions from the CCR5

preparations may also include solutions for intranasal administration. opacifiers for oral solutions, suspensions and emulsions. Liquid form glycol solutions for parenteral injection or addition of sweeteners and emulsions. As an example may be mentioned water or water-propylene Liquid form preparations include solutions, suspensions and

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8 pharmaceutically acceptable carrier, such as an inert compressed gas, e.g. and solids in powder form, which may be in combination with a Aerosol preparations suitable for inhalation may include solutions

 \aleph parenteral administration. Such liquid forms include solutions, suspensions converted, shortly before use, to liquid form preparations for either oral or and emulsions. Also included are solid form preparations which are intended to be

မွ patch of the matrix or reservoir type as are conventional in the art for this lotions, aerosols and/or emulsions and can be included in a transdermal transdermally. The transdermal compositions can take the form of creams The compounds of the invention may also be deliverable

Preferably the compound is administered orally.

႘ၟ In such form, the preparation is subdivided into suitably sized unit doses amount to achieve the desired purpose containing appropriate quantities of the active component, e.g., an effective Preferably, the pharmaceutical preparation is in a unit dosage form.

WO 00/66559

PCT/US00/11633

to the particular application. about 25 mg to about 300 mg, more preferably from about 50 mg to about be varied or adjusted from about 10 mg to about 500 mg, preferably from 250 mg, and most preferably from about 55 mg to about 200 mg, according The quantity of active compound in a unit dose of preparation may

daily dosage may be divided and administered in portions during the day as condition being treated. Determination of the proper dosage regimen for a depending upon the requirements of the patient and the severity of the particular situation is within the skill of the art. For convenience, the total The actual dosage of CCR5 compound employed may be varied

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귥 considering such factors as age, condition and size of the patient as well as preferably about 200 mg/day, in two to four divided doses. dosage regimen for oral administration can range from about 100 mg/day severity of the symptoms being treated. A typical recommended daily compounds of the invention and/or the pharmaceutically acceptable salts to about 300 mg/day, preferably 150 mg/day to 250 mg/day, more thereof will be regulated according to the judgment of the attending clinician The amount and frequency of administration of the CCR5

25 8 other agents used in combination with the CCR5 antagonists will be severity of the condition treated. dosage regimens in the package inserts or as set forth in the protocols, determined by the attending clinician inview of the approved doses and taking into consideration the age, sex and condition of the patient and the The doses and dosage regimens of the NRTIs, NNRTIs, PIs and

present invention by the methodology of Amplicor -1 Monitor 1.5 (available plasma of the patient as measured by quantitative, multi-cycle reverse transcriptase PCR methodology. HIV-1-RNA is preferably measured in the HIV-1-RNA" in the context of the present invention means that there are the HIV-1-RNA viral load below the detectable limit. The "detectable limit of from Roche Diagnsotics) or of Nuclisens HIV-1 QT -1. fewer than about 200 to fewer than about 50 copies of HIV-1-RNA per ml of The goal of the HIV-1 therapy of the present invention is to reduce

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the spirit and scope of the present invention. All such alternatives, modifications and variations are intended to fall within and variations thereof will be apparent to those of ordinary skill in the art. the specific embodiments set forth above, many alternatives, modifications While the present invention has been described in conjunction with

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WHAT IS CLAIMED IS:

A compound represented by the structural formula II

or a pharmaceutically acceptable salt thereof, wherein (1) X^a is $-C(R^{13})_{2^*}$, $-C(R^{13})(R^{19})_{7^*}$, $-C(O)_{7^*}$, $-O_{7^*}$, $-NH_{7^*}$, $-N((C_{7^*}C_{6})alkyl)_{7^*}$

$$Q-C(Q)-N((C_1-C_6)alkyl)_2$$
 $NR^5-C(Q)-(C_1-C_6)alkyl$

Q-C(O)-(C₁-C₆)alkyl Q-C(O)-O-(C₁-C₆)alkyl Q-C(O)-NH-(C₁-C₆)alkyl Q-CR¹³- ,-CR¹³-

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O-C(O)-N((C₁-C₆)alkyl)₂
$$NR^5$$
-C(O)-(C₁-C₆)alkyl $-CR^{13}$

NR⁵-C(O)-O-(C₁-C₆)alkyl NR⁵-C(O)-NH-(C₁-C₆)alkyl

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$$\begin{array}{lll} NR^5\text{-C(O)-N-((C_1\text{-}C_6)alky!)}_2 & \text{C(O)-(C_1\text{-}C_6)alky!} \\ & \text{-CR}^{13} - & \text{or -N-} \end{array};$$

R1 is hydrogen, C1-C6 alkyl or C2-C6 alkenyl; Ra is R6a-phenyl, R6a-pyridyl, R6a-thiophenyl or R6-naphthyl;

8 R¹⁰, R¹¹-substituted 5-membered heteroaryl; naphthyl; fluorenyl; R¹⁷ R¹² R¹⁷ heteroaryl; R7, R8, R9-substituted 6-membered heteroaryl N-oxide: R² is R⁷, R⁸, R⁹-phenyl; R⁷, R⁸, R⁹-substituted 6-membered

diphenylmethyl R¹⁸ – R3 is R10-phenyl, pyridyl, pyrimidyl, pyrazinyl or thiazolyl; -C-heteroaryl

8 -CH2C(O)NH2, -CH2C(O)-NH(C1-C6)alkyl or -CH2C(O)-N((C1-C6)alkyl)2; -CH₂CH₂OH, -CH₂CH₂-O-(C₁-C₆)alkyl, -CH₂C(O)-O-(C₁-C₆)alkyl, R4 is hydrogen, C1-C6 alkyl, fluoro-C1-C6 alkyl, cyclopropylmethyl,

hydrogen and (C₁-C₆)-alkyl; R5 and R11 are independently selected from the group consisting of

consisting of hydrogen, halogen, -CF3, CF3O-, -CN, -CF3SO2-, R12-phenyl R6a is 1 to 3 substituents independently selected from the group

-NHCOCF₃, 5-membered heteroaryl and -N, wherein X is -O-, -NH.

R⁶ is independently selected from the group consisting of R^{6a} and

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(C₁-C₆)alkyl, halogen, -NR[∞]R²¹, -OH, -CF₃, -OCH₃, -O-acyl, and -OCF₃; R9 is R7, hydrogen, phenyl, -NO2, -CN, -CH2F, -CHF2, -CHO, R7 and R8 are independently selected from the group consisting of

5 cycloalkyl, \cdot SR²³, \cdot SOR²³, \cdot SO₂RR²³, \cdot SO₂NH(C₁·C₈ alkyl), \cdot OSO₂(C₁·C₆)alkyl, \cdot OSO₂CF₅, hydroxy(C₁·C₆)alkyl, \cdot CON R²⁰R²¹, \cdot CON(CH₂CH₂·O·C₈)alkyl, \cdot CON(CH₂CH₂·O·C₈)alkyl, \cdot CON R²⁰R²¹, $cycloalkyl(C_1-C_0)alkyl), -NHCO(C_1-C_0)alkyl, -NHCOCF_3, -NHSO_2N((C_1-C_0)alkyl), -NHCOCF_3, -NHSO_2N((C_1-C_0)alkyl)$ -CH=NOR²⁰, pyridyl, pyridyl N-oxide, pyrimidinyl, pyrazinyl, C₆)alkyl)₂, -NHSO₂(C₁-C₆)alkyl, -N(SO₂CF₃)₂, -NHCO₂(C₁-C₆)alkyl, C₃-C₁₀ -N(R²⁰)CONR²¹R²², -NHCONH(chloro-(C₁-C₆)alkyl), -NHCONH((C₃-C₁₀)-

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-OCONH(C₁-C₆)alkyl, -CO₂R²⁰, -Si(CH₃)₃ or -B(OC(CH₃)₂)₂: R¹⁰ is (C₁-C₆)alkyl, -NH₂ or R¹²-phenyl;

consisting of hydrogen, (C₁-C₆) alkyl, -CF₃, -CO₂R₂₀, -CN, (C₁-C₆)alkoxy R12 is1 to 3 substituents independently selected from the group

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consisting of hydrogen and (C₁-C₆)alkyl; R13, R14, R15 and R16 are independently selected from the group

group and with the carbon to which they are attached form a spiro ring of 3 hydrogen and C₁-C₆ alkyl, or R¹⁷ and R¹⁸ together are a C₂-C₅ alkylene to 6 carbon atoms; R17 and R18 are independently selected from the group consisting of

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C10)cycloaikyl(C1-C8)alkyl or (C1-C8)alkoxy(C1-C8)alkyl; R" is R'-phenyl, R'-heteroaryl, R'-naphthyl, C3-C1, cycloalkyl, (C3-

consisting of H and C1-C6 alkyl; and R^{20} , R^{21} and R^{22} are independently selected from the group

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R²³ is C₁-C₈ alkyl or phenyl; or

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Xª is -C(R13)(R19)-, -C(O)-, -O-, -NH-, -N((C1-C6)alkyl)-,

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PCT/US00/11633

WO 00/66559

OR³ CH₂-(C₁-C₅)alkyl-R³ NOR⁴⁸ O-C(O)-(C₁-C₆)alkyl-CR¹³-, -CR¹³-, -

O-C(O)-(C₁-C₆)alkyl O-C(O)-NH-(C₁-C₆)alkyl O-C(13_ , -CR¹³_ ,

O-C(O)-N((C₁-C₆)alkyl)₂ NR^5 -C(O)-(C₁-C₆)alkyl $-CR^{13}$ - $-CR^{13}$ -

NR⁵-C(O)-O-(C₁-C₆)alkyl NR⁵-C(O)-NH-(C₁-C₆)alkyl

 $\begin{array}{lll} NR^5\text{-}C(O)\text{-}N\text{-}((C_1\text{-}C_6)\text{alkyl})_2 & C(O)\text{-}(C_1\text{-}C_6)\text{alkyl} \\ -CR^{13}\text{--} & \text{or }\text{-}N\text{--} \\ \end{array};$

ಕ CH2C(O)-NH-(C1-C6)alkyl or -CH2C(O)-N((C1-C6)alkyl)2; -CH2CH2-O-(C1-C6)alkyl, -CH2C(O)-O-(C1-C6)alkyl, -CH2C(O)NH2, Ra is R6b-phenyl, R6b-pyridyl or R6b-thiophenyl; R^{4a} is fluoro-C₁-C₆ alkyl, cyclopropylmethyl, -CH₂CH₂OH, R6b is CH₃SO₂-; and

5 R1, R2, R3, R5, R14, R15, R16 and R19 are as defined in (1).

The compound of claim 1 wherein Ra is

20 -C(R13)(R18)- or -C(=NOR4)-. The compound of claim 1, formula II(1), wherein Xa is -CHOR3

R" is hydrogen and R" is R'-phenyl. The compound of claim 3 wherein R3 is pyridyl, R4 is (C1-C6)alkyl, or

25 -C(R¹³)(R¹⁸)- or -C(=NOR⁴B)-. The compound of claim 1, formula II(2), wherein Xa is -CHOR3,

cyclopropylmethyl or trifluoroethyl, or R13 is hydrogen and R19 is R6-phenyl The compound of claim 5 wherein R3 is pyridyl, R4a is

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R9-pyridyl or an N-oxide thereof; or R7, R8, R9-pyrimidyl. The compound of claim 1 wherein R2 is R7, R8, R9-phenyl; R7, R8,

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8. The compound of claim 7 wherein R² is selected from the group

wherein R7 and R8 are selected from the group consisting of (C1-C6)alkyl, halogen, and -NH₂, and R⁹ is hydrogen.

represented by the formula A compound of claim 1 selected from the group consisting of those

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| F ₃ CO- | F ₃ CO- | нзсsо ₂ - | H ₃ CSO ₂ - | F ₃ C- | H ₃ CSO ₂ - | H3CSO2- | B | Br | Br | Br | CH3SO2- | Br | Br | Br |
| -0 -0 -2 | CH. N. | -CH. | _ z ∕_> | + × | N N N N N N N N N N N N N N N N N N N | _ ₹_> Ç> | ÷-0 | [‡] → × | Ş> Z-> | Š | - o- N | | | - OF N |
| CH ₂ | н₃с Сн₃ | H ₃ C, CH ₃ | H₃C CH₃ | н₃СДСН3 | сн₃⋛он | CI NH ₂ | H ₃ C NH ₂ | H ₃ C _N CH ₃ | сн, Дон | CI NH ₂ | H ₃ C CH ₃ | н₃С₩СН₃ | н₃С∰СН₃ | н₃С-Ж сн₃ |

- 84 -

| F ₃ CO- | F ₃ C- | н ₃ СSО ₂ - | H3CSO2- | H3CSO2- | H3CSO2- | H ₃ CSO ₂ - | H ₃ CSO ₂ - | Br | Br | F ₃ CO- | Br | Br | F ₃ CO- |
|---------------------------------------|--|-----------------------------------|----------------------------------|---------|-----------|-----------------------------------|---------------------------------------|---|---|--------------------|--------------------|----------|---|
| 5-0 | ÷- | S CH | | S OH- | CH. N. N. | -0 -0 -0 -0 -0 | r P | C. O. N. N. | -CH. | l o v | 0-0 2-0 | | |
| H ₃ C N CH ₃ | H ₃ C _C H ₃ | н эс Снэ | H ₃ C CH ₃ | н₃сДсн₃ | нас Сна | н₃СДСН₃ | H ₃ C N CH ₃ | H ₃ C ₁ CH ₃ | H ₃ C ₁ CH ₃ | #5 7-043 | PH CHANGE | Н3С, СН3 | H ₃ C _N CH ₃ |

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| 711 | 711 | 711 | F ₃ C- | H3CSO2- | H3CSO2- | F ₃ C- | H3CSO2- | Br | Ω | Ω | Ω | Ω | F3CO- |
| -ch. | H ON N | | - Ç₽ - OP | | -0 -0 -0 -2 -2 | [†] → | -04- S-04- | ÷-0 | - <u>C</u> | ÷- | ţ, Ç | ÷-0 | ن پ |
| H ₃ C CH ₃ | H ₃ C ₂ CH ₃ | H ₃ C _N CH ₃ | CC N | H ₃ C CH ₃ | | H ₃ C N N N CH ₃ | H ₃ C CH ₃ | CI N | H ₃ C N N N N | CI N | H ₃ C CH ₃ | н₃С, Сн₃ | |

- 85 -

PCT/US00/11633

- 86 -

| | Вг | Br | п | F ₃ C- | F ₃ C- | F ₃ C- | Br | ₿r | Br | Вг | | Ω |
|--------------|---------------------------------------|----------------------------------|---|---------------------------------------|-----------------------------------|-------------------|-----|--------------------------------------|-----|---|----------|-------------|
| Enantiomer A | N. | -с- | -c+- | Ť. Č. | Ť-o | CH. N | | H ₃ CH Q N | | - 0 H | H N | ÷ >zv |
| | H ₃ C N CH ₃ | н,с, ₩ сн, | CH. V | H ₃ C N CH ₃ | H ₃ C, CH ₃ | Ω Ω Ω | 0-z | H ₃ C NCH ₃ | H3C | H ₃ C _N CH ₃ | O+z \ CI | Ω Ω Ω |

| | Ω | Ō | F ₃ CO- | F ₃ CO- | F ₃ CO- | F ₃ CO- | F ₃ CO- | F3CO- | Br | Br | Br |
|--------------|---|---|----------------------------------|---------------------------|----------------------------------|--------------------|----------------------------------|--|--|--|---------------------------------|
| Enantiomer A | , | -CH- Enantiomer A | —CH- Enantiomer B | N -CH- Enantiomer A | CH- CH- Enantiomer A | N O | O V N Y | O N N Enantiomer A | - CH- | -CH- | CH- Enantiomer A |
| 요- 3 | H ₃ C ₁ CH ₃ | H ₃ C _N CH ₃ | H ₃ C CH ₃ | H3C N N N OCH3 | H ₃ C CH ₃ | Н3С СН3 | H ₃ C CH ₃ | H ₃ C H ₃ C H ₃ C | H ₃ C _Y C _Y | H ₃ C N CH ₃ | H ₃ CCH ₃ |

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| | CH ₃ CH ₂ Q N | H3CO_N | H ₃ CO/N | - C= NOCH3 | - C-N OCH3 | сн ₃ сн ₂ Q N —с— | NOCH3 | NOCH3 | E-isomer | H ₃ CO_N — C — Z-isomer | O N N N N N N N N N N N N N N N N N N N | -CH- Enantiomer B | -CH- Enantiomer B |
| • | н₃с ₹он | H ₃ C NH ₂ | н₃с√тон | H ₃ C\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\ | CI NH2 | н₃с, Сн₃ | CI NH ₂ | H ₃ C NH ₂ | н _з с Сн _з | н _з с Сн _з | H ₃ C ₇ CH ₃ N ₇ N NH ₂ | H ₃ C ₁ CH ₃ N CH ₃ | N N |

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| | B | B | F,C. | Br | Br | Br | Br | Br | Br | Br | . Br | Br | Br |
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| | CH ₃ CH ₂ O- _N -C- | CH ₃ CH ₂ O _{>N} -C- | CH ₃ CH ₂ O-N -C- | CH ₃ CH ₂ O-N -C- | CH ₃ CH ₂ O-N -C- | CH ₃ CH ₂ Q N C | 13CO | H. J. | -≥^0 | сн ₃ сн ₂ Q N —с— | H3C > | CH3CH2Q N -C- | CH ₃ CH ₂ Q -C- |
| 3013 | H ₃ C N ₇ N SCH SCH SCH SCH SCH SCH SCH SCH SCH SCH | N.O CH ₃ | H ₃ C ₁ CH ₃ N≪N | H ₃ C N N CH ₃ CH ₃ | O+Z Br | | н₃С√тсн₃ | H ₃ C CH ₃ | H ₃ C N CH ₃ | Br. Br | H ₃ C CH ₃ | H ₃ C ₁ CH ₃ OH | O 4-X CH ₃ |

| H ₂ C CH ₂ C CH ₂ C CH ₂ C CH ₃ C CH | ١ | | | | L | | | | |
|---|------|---|---|-----|---|--------|-------------------|-------------|----------------------|
| | | CH ₃ CH ₂ O ₂ N -C- | CH ₃ CH ₂ O _{>N} -C- | | | , | CH3CH2Q N C | -с- -с- | H ₃ C O N |
| | 3000 | ν-ζ-γ- γ-ζ-γ-ξ- γ-ζ-γ-ξ- | CH ₃ | ž 🗸 | 모 | O+N Br | | z√ } | Z-{-}- |

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PCT/US00/11633

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| F3CO- | F3CO- | F ₃ C- | F ₃ C- | F3CO- | F ₃ CO- | F ₃ C- | F ₃ CO- | F3CO- | F ₃ CO- | F ₃ C- | F ₃ C- | F3C0- | F ₃ C. | F ₃ C- |
|----------------------------------|---|--|--|--|---|--|--|----------|---|--|---------------------------------------|----------------------|----------------------------------|--------------------|
| | | Y | , Y | Р | Ŷ | Y | P | Ŷ | P | · · | γ, | Ϋ́ | δ | · |
| H3CO(CH2)2O N -C- | CH ₃ O _N N E isomer | сн _з сн ₂ о, N —с- | СН ₃ СН ₂ О N С- | CH ₃ CH ₂ O, N -C- | сн ₃ о_ N | CH ₃ O _N N —C— E isomer | CH ₃ CH ₂ O, N -C- | <u>م</u> | CH ₃ CH ₂ Q N -C- | CH ₃ CH ₂ O, N -C- | CH ₃ O ₋ N N | CH3CH2O, N -C- | | - C- N |
| H ₃ C CH ₃ | H ₃ C CH ₃ | # # # | #5 ₹-64 #6 | H3C 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 | #c Act Act Act Act Act Act Act Ac | ₽° C | ну С | нзс Снз | HC Con | Но ₹ | #c ₩c# | CI NH ₂ | H ₃ C NH ₂ | CI NH ₂ |

| F ₃ C- | Br | F3CO- | F3CO- | F ₃ CO- | F3CO- | F3CO- | F ₃ CO- | F3C0- | F ₃ CO- | F3CO- | F3CO- | F3CO- | F ₃ CO- |
|----------------------------------|---------------------------|---|--|--|---|---------------------|---|---------------------|--------------------|---|------------|----------|---|
| CH3CH2Q N —C— | CH ₃ O, -C- | СН ₃ (СН ₂₎₂ О, N —С- | CH ₃ (CH ₂) ₂ O ₄ N -C- | CH ₃ (CH ₂) ₂ Q, N -C- | CH ₃ CH ₂ O, N | - C- - C- - N | CH3CH2O, N | CH3CH2Q N -C- | CH ₃ O, | сн ₃ сн ₂ о, N | ٢ اب=×٥ | 4=20 | CH ₃ CH ₂ O, N |
| H ₃ C CH ₃ | CI NH ₂ | 40 A CH | 0Z | H ₃ C N N N N | 0+z | 0 | H ₃ C N N N N N | CI N | CI N CI | CI N | н₃С्Хон | Hc Ly ch | HC CHO |

- 96 -

| п | 71) | 711 | Br | 뫈 | CH3SO2- | Br | Br | Br | Br | Br | Br | Br |
|-----|-------------------------------------|----------------------------------|--|----------------------------------|---------|---|--------------------|--------------------|-------------------------|----------------------------------|----------------------|--------------------|
| | t 🏷 | -t | -c | - T ~~ | ţ-° | | -CH ₂ - | -CH ₂ - | -CH ₂ - | -CH ₂ - | -CH ₂ - | -CH ₂ - |
| F G | H ₃ C CH ₃ | H ₃ C CH ₃ | H ₃ C N N N CH ₃ | H ₃ C CH ₃ | 45 A-CH | H ₃ C N N N N CH ₃ | ~ | | H ₃ C O-N Cl | H ₃ C-NH ₂ | H ₃ C - N | H ₃ C C |

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|---|---|--------------------|----------------------------------|---|---|---------------------------------------|----------------------------------|----------------------------------|--|--------------------------------------|--|--------------------------------------|---|
| | F,C- | F ₃ CO- | F3CO- | F ₃ C- | CH3SO2- | СН3SO2- | F3CO- | F3CO- | CH3SO2- | сн ₃ ѕо ₂ . | F ₃ C- | Ω | Br |
| | cı Do | -HC- | -μc- | | - - | † <u> </u> | | ţ-o | | ¦₀ † ≿> | F ≿ | t S | t op |
| | H ₃ C ₁ CH ₃ | Hoc Hoch | H ₃ C CH ₃ | H ₃ C N N N N N | H ₃ C N N N N N | H ₃ C N CH ₃ | H ₃ C CH ₃ | H ₃ C CH ₃ | H ₃ C ₁ CH ₃ N N N | H ₃ C N N N N | to the state of th | H ₃ C N N N N | H ₃ C√CH ₃ N≪N |

- 97 -

F3C0 F₃CO F3C0-F3CO-I £ Enantiomer II Enantiomer II Enantiomer II

10. A compound selected from the group consisting of

effective amount of a CCR5 antagonist of claim 1 in combination with a dermatitis, psoriasis, asthma, allergies or multiple sclerosis, comprising an disease, arthritis, rheumatoid arthritis, inflammatory bowel disease, atopic Immunodeficiency Virus, solid organ transplant rejection, graft v. host A pharmaceutical composition for the treatment of Human

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pharmaceutically acceptable carrier.

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WO 00/66559

PCT/US00/11633

- 98 -

- or multiple sclerosis. inflammatory bowel disease, atopic dermatitis, psoriasis, asthma, allergies transplant rejection, graft v. host disease, arthritis, rheumatoid arthritis, medicament for treating Human Immunodeficiency Virus, solid organ The use of a compound of claim 1 for the preparation of a
- useful in the treatment of Human Immunodeficiency Virus, medicament for combined use with one or more antiviral or other agents The use of a compound of claim 1 for the preparation of a

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- group consisting of nucleoside reverse transcriptase inhibitors, nonnucleoside reverse transcriptase inhibitors and protease inhibitors. The use of claim 13 wherein the antiviral agent is selected from the
- 5 organ transplant rejection, graft v. host disease, inflammatory bowel 5 disease, rheumatoid arthritis or multiple sclerosis. medicament for combined use with one or more agents for treating solid The use of a compound of claim 1 for the preparation of a
- 25 20 structural formula t: or multiple sclerosis, wherein the CCR5 antagonist is represented by the inflammatory bowel disease, atopic dermatitis, psoriasis, asthma, allergies transplant rejection, graft v. host disease, arthritis, rheumatoid arthritis, medicament for treating Human Immunodeficiency Virus, solid organ The use of a CCR5 antagonist of formula I for the preparation of a

or a pharmaceutically acceptable salt thereof, wherein X is $-C(R^{13})_{2^-}$, $-C(R^{13})(R^{19})_{1^+}$, $-C(O)_{1^+}$, $-O_{1^+}$, $-N((C_{1^+}C_{6})alky)_{1^+}$,

NR⁵-C(O)-N-((C₁-C₆)alkyl)₂ C(O)-(C₁-C₆)alkyl | or -N-

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R¹ is hydrogen, C₁-C₆ alkyl or C₂-C₆ alkenyl; R is R6-phenyl, R6-pyridyl, R6-thiophenyl or R6-naphthyl;

heteroaryl; R7, R8, R9-substituted 6-membered heteroaryl N-oxide; R2 is R7, R8, R9-phenyl; R7, R8, R9-substituted 6-membered

R10, R11-substituted 5-membered heteroaryl; naphthyl; fluorenyl; R17 _____R12 ___R17 diphenylmethyl -C-heteroaryl or R¹⁸

-CH2CH2OH, -CH2CH2-O-(C1-C6)alkyl, -CH2C(O)-O-(C1-C6)alkyl, R⁴ is hydrogen, C₁-C₆ alkyl, fluoro-C₁-C₆ alkyl, cyclopropylmethyl, R3 is R6-phenyl, R6-heteroaryl or R6-naphthyl;

5 -CH₂C(O)NH₂, -CH₂C(O)-NH(C₁-C₆)alkyl or -CH₂C(O)-N((C₁-C₆)alkyl)₂; hydrogen and (C₁-C₆)-alkyl; R5 and R11 are independently selected from the group consisting of

CH₃C(O)-, -CN, CH₃SO₂-, CF₃SO₂-, R¹⁴-phenyl, R¹⁴-benzyl, consisting of hydrogen, halogen, C₁-C₆ alkyl, C₁-C₆ alkoxy, -CF₃, CF₃O-R6 is 1 to 3 substituents independently selected from the group

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 $\label{eq:ch3c} \text{CH}_3\text{C}(=\text{NOCH}_3\text{P}, \text{ CH}_3\text{C}(=\text{NOCH}_2\text{CH}_3\text{P}, \text{ O})}, \text{CH}_3\text{C}(=\text{NOCH}_3, \text{-NH}_2, \text{-NHCOCF}_3, \text{-NH}_2, \text{-NHCOCF}_3, \text{-NH}_2, \text{-NHCOCF}_3, \text{-NHCOCF}$ $NHCONH(C_1-C_6 alkyl)$, $-NHCO(C_1-C_6 alkyl)$, $-NHSO_2(C_1-C_6 alkyl)$

25 (C₁-C₆)alkyl, halogen, -NR²⁰R²¹, -OH, -CF₃, -OCH₃, -O-acyl, and -OCF₃; 5-membered heteroaryl and $\stackrel{-N}{\searrow}_x$, wherein X is -O-, -NH- or $-N(CH_3)$ -; $\ensuremath{\mbox{R}^{7}}$ and $\ensuremath{\mbox{R}^{8}}$ are independently selected from the group consisting of

 $C_0)alkyl)_2, \ -NHSO_2(C_1-C_0)alkyl, \ -N(SO_2CF_3)_2, \ -NHCO_2(C_1-C_0)alkyl, \ C_3-C_1)alkyl, \ C_3-C_2(C_1-C_0)alkyl, \ C_3-C_2(C_1-C_0)al$ ${\it cycloalkyl}(C, -C_0) alkyl), -{\it NHCO}(C, -C_0) alkyl, -{\it NHCOCE}_3, -{\it NHSO}_2 N((C, -C_0) alkyl), -{\it NHCO}_2 N((C, -C_0) alkyl), -{\it NHCO}_2 N((C, -C_0) alkyl), -{\it NHCOCE}_3, -{\it NHSO}_2 N((C, -C_0) alkyl), -{\it NHSO}_2$ -N(R^∞)CON R^2 ' R^{22} , -NHCONH(chloro-(C,-C₆)alkyl), -NHCONH((C₃-C₁₀) -CH=NOR[∞], pyridyl, pyridyl N-oxide, pyrimidinyl, pyrazinyl, R9 is R7, hydrogen, phenyl, -NO2, -CN, -CH2F, -CHF2, -CHO,

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WO 00/66559

PCT/US00/11633

cycloalkyl, -SR23, -SOR23, -SO2R23, -SO2NH(C,-Ce alkyl), -OSO2(C,-Ce)alkyl, -OCONH(C,-C₆)alkyl, -CO₂R $^{\infty}$, -Si(CH₃)₃ or -B(OC(CH₃)₂)₂; -OSO₂CF₃, hydroxy(C₁-C₆)alkyl, -CON R²⁰R²¹, -CON(CH₂CH₂-O-CH₃)₂, R¹⁰ is (C₁-C₆)alkyl, -NH₂ or R¹²-phenyl;

consisting of hydrogen, (C₁-C₆) alkyl, -CF₃, -CO₂R₂₀, -CN, (C₁-C₈)alkoxy R¹² is 1 to 3 substituents independently selected from the group

consisting of hydrogen and (C₁-C₆)alkyl; R13, R14, R15 and R16 are independently selected from the group

5 group and with the carbon to which they are attached form a spiro ring of 3 hydrogen and C1-C6 alkyl, or R17 and R18 together are a C2-C5 alkylene to 6 carbon atoms; R¹⁷ and R¹⁸ are independently selected from the group consisting of

C₁₀)cycloalkyl(C₁-C₈)alkyl or (C₁-C₈)alkoxy(C₁-C₈)alkyl; R¹⁹ is R^e-phenyl, R^e-heteroaryl, R^e-naphthyl, C₃-C₁₀ cycloalkyl, (C₃- R^{∞} , R^{21} and R^{22} are independently selected from the group

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consisting of H and C,-C₆ alkyl; and R²³ is C₁-C₆ alkyl or phenyl.

8 **1**7 The use of claim 16 wherein R is $\mathbb{R}^6 - \{\}$

-C(=NOR4)-. 18. The use of claim 16 wherein X is -CHOR3, -C(R13)(R19)- or

25 is hydrogen and R16 is R6-phenyl. The use of claim 18 wherein R3 is pyridyl, R4 is (C1-C6)alkyl, or R13

pyridyl or an N-oxide thereof, or R7, R8, R9-pyrimidyl. The use of claim 16 wherein R² is R⁷, R⁸, R⁹-phenyl, R⁷, R⁸, R⁹.

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The use of claim 20 wherein R2 is selected from the group

$$R^{7}$$
 R^{8}
 R^{7}
 R_{9}
 R_{9}

WO 00/66559 PCT/US00/11633

halogen, and -NH $_2$, and R 9 is hydrogen. wherein ${\sf R}^7$ and ${\sf R}^8$ are selected from the group consisting of (C1-C6)alkyl.

treatment of Human Immunodeficiency Virus. Virus, further comprising one or more antiviral or other agents useful in the The use of claim 16 for the treatment of Human Immunodeficiency

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- group consisting of nucleoside reverse transcriptase inhibitors, non-The use of claim 22 wherein the antiviral agent is selected from the
- **5** nucleoside reverse transcriptase inhibitors and protease inhibitors.
- useful in the treatment of said diseases. arthritis or multiple sclerosis, further comprising one or more other agents rejection, graft v. host disease, inflammatory bowel disease, rheumatoid The use of claim 16 for the treatment of solid organ transplant

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pharmaceutical composition comprising an effective amount of a CCR5 Human Immunodeficiency Virus in a pharmaceutically acceptable carrier. an effective amount of a antiviral or other agent useful in the treatment of separate containers, one or more pharmaceutical composition comprising antagonist of claim 16 in a pharmaceutically acceptable carrier, and in pharmaceutical compositions for use in combination to treat Human Immunodeficiency Virus which comprises in one container a A kit comprising in separate containers in a single package

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INTERNATIONAL SEARCH REPORT

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| neame and meaning address of the ISA. European Parent Office, P.B. 5918 Peterotean 2 NI. – 2250 NF Afjandy. Tel. (431–70) 340–3010 Tz. 31 651 spo nt. Faut (431–70) 340–3010 Tz. 31 | August 2000 | Date of the ectual completion of the international search | material material reletor | Purper documents are stand in the continuation of box C. Special categories of cited documents : | | WO 98 01425 A (SCHERING CORP) 15 January 1998 (1998-01-15) claims 1,11; examples | WO 98 06697 A (SCHERING CORP) 19 February 1998 (1998-02-19) claims 1,11; examples | WO 98 05292 A (SCHERING CORP) 12 February 1998 (1998-02-12) claims 1,13; examples & US 5 889 006 A 30 March 1999 (1999-03-30) cited in the application | Clation of document, with indication, where appropriate, of the relevant passages | C. DOCUMENTS CONSIDERED TO BE RELEVANT | Boardario dala base consulted during the international search (name of data base and, where practical, search name used) EPO-Internal, CHEM ABS Data, WPI Data | IPC 7 CO7D A61K A61P Commentation searched other than minimum documentees on to the extent that each documents are included in the fields searched | A wing to international Patent Classification (IPC) or to both national classification and IPC B. RELDS SEARCHED Minimum documentation searched (classification persons information by the classification by | TPC 7 CONDUCTIVE MATER A61K31/4523 A61P31/12 A61P19/00 | |
| Authorized officer De Jong, 8 | 11/08/2000 | Date of mailing of the international search report | I also gouthest published after the framidous fling data printly data and not in condition with the uppsication but cited to indeterand the principle or theory underlying the framidon. The principle of theory underlying the Y. document of purificate relevance; the clitical interaction certain to considered royal or carrior be considered to hirdle an inventive stap when the document is taken used the principle of the principle of the principle of the principle of constant is considered to limited the principle also when document is considered with one or more other was the mental card combination being dockdas to a person desired in the art. | X Patent family members are listed in arreax | -/- | | | | relevant passages | | base and, where practical, exects terms use | caum epinous) at each documents are included in the fields. | Affication and IPC | C07D413/14 | PCT/US 00/11633 |
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C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT
CARGOY Citation of document, with midestorum are appropriate, of the relevant passages WO 99 04794 A (OATES BRYAN :FINKE PAUL E ¿ (US): MACCOSS MALCOLM (US): MERCK & CO I) "4 February 1999 (1999-02-04) abstract: claim 1 PCT/US 00/11633 Relevant to ctom No. 1,16

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INTÉRNATIONAL SEARCH REPORT

Inter and Application No

WO 9904794 WO 9801425 Patent document cited in search report W0 9806697 WO 9805292 Þ > 04-02-1999 15-01-1998 19-02-1998 12-02-1998 Publication AU 3973297 AC 1232453 AC 1232453 AC 9900433 AC 990052029 AC 1P 2000500786 T NO 990671 AC 931536 AC 1231536 AC 123156 AC 독무요은 SARRORFSAR S 5889006 | S 5889007 | S 5889007 | S 5889007 | S 589907 | S 589907 | S 589007 | S 58900 Patent family member(s) 3581097 2259655 0912515 11514671 8576098 1003514 02-02-1998 15-01-1998 06-05-1999 14-12-1999 16-02-1999 31-05-2000 06-03-1998 20-10-1999 14-07-1999 16-06-1999 25-01-2000 15-04-1999 19-07-1999 30-03-1999 25-02-1998 23-11-1999 20-110-1999 16-06-1999 01-09-1999 02-02-2000 07-04-1999 19-07-1999 28-03-2000 Publication date

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